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(54) Title: COMPOSITIONS FOR USE IN IDENTIFICATION OF MIXED POPULATIONS OF BIOAGENTS

(57) Abstract: The present invention provides oligonucleotide primers, compositions, and kits containing the same for rapid identification of bacterial bioagents and populations of bioagents which are members of the Staphylococcus bacterial genus by amplification of a segment of bioagent nucleic acid followed by molecular mass analysis.

COMPOSITIONS FOR USE IN IDENTIFICATION OF MIXED POPULATIONS OF BIOAGENTS

CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] This application claims the benefit of priority to U.S. Provisional Application Serial No. 60/896,801, filed March 23, 2007, the disclosure of which is incorporated by reference in its entirety for any purpose.

SEQUENCE LISTING

[0002] Computer-readable forms of the sequence listing, on CD-ROM, containing the file named DIBIS0093WOSEQ.txt, which is 69,632 bytes (measured in MS-DOS), and were created on March 22, 2007, are herein incorporated by reference.

STATEMENT OF GOVERNMENT SUPPORT

[0003] This invention was made with support from NIH/NIAID, contract: 1 UC1-AI067232-01, project: 842. The U.S. government has certain rights in the invention.

FIELD OF THE INVENTION

[0004] The present invention relates generally to the field of genetic identification and quantification of bioagents, including mixed populations of bioagents and provides methods, compositions and kits useful for this purpose, as well as others, when combined with molecular mass analysis.

BACKGROUND OF THE INVENTION

[0005] Drug resistance is a growing problem in disease treatment and control. Development of antibiotic resistance by bacteria, especially to broad-range antibiotics, is particularly problematic.

Resistance emerges as use and/or misuse of drugs provides a selection advantage to resistant populations of infectious bioagents. Effective surveillance of emerging drug resistance is important for identifying, monitoring and controlling resistant populations and for developing appropriate treatment strategies.

[0006] Use of drugs to treat infection with bioagents having a propensity towards resistance can lead to treatment failure and/or development of new drug resistance. Furthermore, the methods available for detection of drug resistance can be prohibitively time consuming and often do not provide sufficient sensitivity or precision to detect low percentages of emerging resistant populations of bioagents. Thus, treatment of patients with certain drugs is often avoided, sometimes resulting in over-use of alternative drugs, and/or development of new drug-resistant strains.

[0007] Quinolones, specifically fluoroquinolones, are highly potent broad-spectrum antibiotics that are used to treat several types of bacterial infections. Because of their widespread use, resistance to quinolones has become prevalent among several classes of bacterial bioagents. A SNP (single-nucleotide polymorphism) within the quinolone resistance determining region (QRDR) of the gyrA gene confers quinolone resistance to Staphylococcus aureus bacteria. Ciprofloxacin, levofloxacin, moxifloxacin and gatifloxacin, among the fluoroquinolones used in treating certain types of Staphylococcus aureus infections, are being used less frequently in certain types of infections due to the risk of drug-resistance development. Methicillin-resistant Staphylococcus aureus (MRSA) strains are particularly adept at developing quinolone resistance, and are thus not typically treated with quinolones. However, the number of antibiotics available for treating bacteria that are resistant to both methicillin and quinolones is limited. Development of sensitive, rapid methods that would enable early detection of quinolone resistant bacteria might allow for the use of quinolones before resistance emerges.

[0008] Standard methods for determining bacterial drug resistance rely on phenotypic characterization. These methods typically require culturing bacteria from a clinical sample for a period of at least 24-48 hours and subsequent susceptibility testing of the cultured bacteria using assays such as agar/broth dilution and/or disk diffusion, which can require an additional 18-24

hours. These tests are relatively insensitive as they rely on visible phenotypic readouts such as culture growth and can only detect a resistant population if it represents a sufficiently high proportion of total bacteria in the sample. Thus, these standard methods are labor intensive, time-consuming, and insensitive, often resulting in misdiagnosis or delay of diagnosis, and by extension, use of inappropriate drug regimens. Thus, there is a long-felt and unmet need for methods that can rapidly detect emerging populations of bioagents and provide sufficient sensitivity and resolution to identify a bioagent that represents only a small percentage of a sample. Specifically, there is a need for methods that can identify small drug-resistant populations in early stages as they emerges in a mixed-population of bioagents, for example, in a sample from a patient being treated with the drug. Such methods would enable monitoring of emerging drug resistance and subsequent design of specific therapeutic approaches tailored to specific bioagent genotypes, and would also reduce the potential for treatment failure and new drug resistance.

SUMMARY OF THE INVENTION

[0009] Provided herein are, inter alia, pairs of primers and compositions comprising pairs of primers; kits comprising the same; and methods for their use in identification of bioagents, populations of bioagents, populations of bioagents, populations of bioagents. The forward and reverse primer members of the pairs of primers are configured to amplify nucleic acids from bioagents, thereby generating amplicons for the nucleic acids. In one aspect, the bioagents are comprised within a population of bioagents. In a preferred embodiment, the primer pairs are configured to amplify one or more nucleic acids from each of the bioagents in the population of bioagents. In one embodiment the primers generate bioagent identifying nucleic acid amplicons. The amplicons are preferably generated from portions of nucleic acid sequences that encode genes essential to antibiotic sensitivity and resistance.

[0010] The primer pairs each comprise a forward and a reverse primer member. In one embodiment, the primer pair is configured to generate an amplicon from within a region defined by SEQ ID NO.: 10, a region of GenBank gi number 49484912, the QRDR (quinolone resistance determining region) of the gyrA gene within this GenBank gi number. In one aspect, either or both of the primer pair members comprise 20 to 35 nucleobases in length. In one aspect the forward

primer pair member comprises at least 70%, at least 80%, at least 90%, at least 95%, at least 97%, or 100% identity to a first portion of SEQ ID NO.: 10. In another aspect, the reverse primer pair member comprises at least 70%, at least 80%, at least 90%, at least 95%, at least 97%, or 100% reverse complementarity to a second portion of SEQ ID NO .: 10. In another embodiment, the forward primer pair member comprises at least 70%, at least 80%, at least 90%, at least 95%, at least 97%, or 100% identity with a portion of SEQ ID NO.: 11, which is a forward primer hybridization region within SEO ID NO., 10. In another embodiment, the reverse primer pair member comprises at least 70%, at least 80%, at least 90%, at least 95%, at least 97%, or 100% reverse complementarity with a portion of SEO ID NO.: 12, a reverse primer hybridization region within SEO ID NO.: 10. In another aspect, the primer pair members are configured to hybridize with at least 70%, at least 80%, at least 90%, at least 95%, at least 97%, or 100% complementarity within a sequence region of a biogent nucleic acid sequence. In one aspect the bioagent nucleic acid sequence is GenBank gi number 49484912. In another aspect, the bioagent nucleic acid sequence is GenBank gi number 57650036. In another aspect, the bioagent nucleic acid sequence is GenBank gi number 47118324. In another aspect, the bioagent nucleic acid sequence is GenBank gi number 27314460

[0011] In one embodiment, the forward primer pair member comprises SEQ ID NO..2 with 0-8 nucleobase deletions, additions and/or substitutions. In another embodiment, the forward primer pair member comprises SEQ ID NO..3 with 0-8 nucleobase deletions, additions and/or substitutions. In another embodiment, the forward primer pair member comprises SEQ ID NO..4 with 0-8 nucleobase deletions, additions and/or substitutions. In another embodiment, the reverse primer pair member comprises SEQ ID NO..5 with 0-6 nucleobase deletions, additions and/or substitutions. In another embodiment, the reverse primer pair member comprises SEQ ID NO..6 with 0-8 nucleobase deletions, additions and/or substitutions. In another embodiment, the reverse primer pair member comprises SEQ ID NO..6 with 0-8 nucleobase deletions, additions and/or substitutions.

[0012] In one embodiment, either or both of the primer pair members comprises at least one modified nucleobase. In one aspect the modified nucleobase is a mass modified nucleobase. In one aspect, the mass modified nucleobase is 5-todo-C. In another aspect the modified nucleobase is a

universal nucleobase. In one aspect, the universal nucleobase is inosine. In another embodiment, either or both of the primer pair members comprise a non-templated 5' T-residue.

[0013] Compositions comprising one or more of the primer pairs and the kits comprising the same, also provided herein, are configured to provide genotyping information, including identification of population genotypes of samples, populations of bioagents, including mixed populations of bioagents.

[0014] Also provided herein are methods of identifying one or more bioagents using the primer pairs and/or kits or compositions comprising the same provided herein.

In one embodiment, the methods are performed for identifying a population genotype for a population of bioagents comprised in the sample. In a preferred embodiment, the population of bioagents is a population of bacterial bioagents. In one embodiment, the population of bioagents comprises two or more bioagents from the same genus, the same species, or even the same strain. In one aspect, the two or more bioagents have the same genotype for one or more locus, gene or nucleotide position. In one embodiment, the population of bioagents is a mixed population of bioagents. In this embodiment, two or more of the bioagents in the population are distinguishable based on one or more characteristics. In one example, the two or more bioagents are distinguishable based on two or more distinct genotypes for a gene, locus, or nucleotide position. In one aspect, the distinct genotype confers resistance to one or more drugs or therapeutic agents. In another aspect, the distinct genotype confers sensitivity to one or more drugs or therapeutic agents. In one embodiment, the mixed population of bioagents comprises a plurality of members of the Staphylococcus genus. In a further embodiment, the population of bioagents comprises a plurality of members of the species Staphylococcus aureus. In one embodiment, the population of bioagents comprises a population of bioagents with two or more distinguishable genotypes for a gene that can confer drug resistance or sensitivity. More preferably, the two or more distinguishable genotypes comprise one genotype that confers resistance to quinolones and another genotype that confers sensitivity to quinolones. In a preferred embodiment, the gene that can confer drug resistance is Gyr A. In a preferred aspect, a distinguishable genotype comprises a C → T transition at nucleotide

within the Gyr A gene, thereby conferring a leucine in place of a serine for the encoded gyrase protein. In a preferred embodiment, the C → T transition is at nucleotide 251 of a sequence extraction with coordinates 7005-9668 (SEQ ID NO.: 8) of GenBank gi number: 49484912, which comprises a nucleotide sequence encoding Gyr A. In one aspect, one or more genotypes is an emerging genotype. In one aspect, the genotype confers drug resistance. In a preferred aspect, the genotype comprises a genotype of the gyrA gene sequence. In one aspect, the genotype comprises a single nucleotide polymorphism.

[0016] In one embodiment, the primer pair is preferably configured to generate an amplicon between about 45 and about 200, more preferably, between about 45 and about 192 linked nucleotides in length within at least a portion of the QRDR region (SEQ ID NO.:10) of the Staphylococcus aureus gyrA gene, which confers quinolone resistance or sensitivity. This region comprises the position of the C → T drug resistance-conferring SNP at within the gyrA gene sequence. The SNP, comprising a change of a single "C" nucleobase to a "T" nucleobase, results in a leucine instead of a serine at amino acid position 84 of the protein. In one aspect, the forward primer is configured to comprise sequence identity within SEQ ID NO.: 11, a region of GenBank gi number 49484912, and the reverse primer is configured to comprise reverse complementarity within SEQ ID NO.: 12, another region of GenBank gi number 49484912. The gyrA primer pairs provided herein, when used in the methods provided herein, can detect a single nucleotide change at this SNP position, and are thus able to determine the drug resistant/sensitive genotype for the gyrA gene for a given Staphylococcus aureus bioagent.

[0017] In one embodiment, the method is performed on a sample that comprises or is suspected of comprising a bioagent or a population of bioagents. In this embodiment, the method comprises obtaining a sample and amplifying a nucleic acid from each of two or more bioagents in the sample using a primer pair provided herein, thereby generating amplicons from the nucleic acids and determining a molecular mass for each of the amplicons using a mass spectrometer. In a preferred embodiment, the determining using a mass spectrometer is accomplished by electrospray ionization mass spectrometry (ESI-MS). In one aspect, the ESI-MS is Fourier transform ion cyclotron resonance mass spectrometry (FT-ICR-MS). In another aspect, it is time of flight (TOF) mass

spectrometry. In another preferred embodiment, the method further comprises calculating a base composition from each molecular mass measurement. In a preferred embodiment, the method further comprises identifying a population genotype for the population of bioagents by comparing each of the molecular mass measurements and/or each of the base compositions calculated from the molecular mass measurements to a database of base compositions and/or molecular masses indexed to the primer pair used in the method and a known bioagent genotype. The database comprises indexed information comprising the molecular mass and/or base composition data that would be derived from a known bioagent having a certain genotype were an amplicon to be generated using the same primer pairs used to amplify nucleic acids in the sample. A match between the experimentally obtained molecular mass and/or base composition obtained by the methods provided herein, for example, on a sample, and a molecular mass and/or base composition comprised in the database correlates a bioagent in the sample with the known bioagent in the database to which the molecular mass and/or base composition is indexed, thus identifying a genotype of that bioagent in the sample. Thus, a sample comprising a population of bioagents that comprises two or more genotypes for the gene or nucleic acid sequence that the primer pair is configured to amplify will correlate with two or more known bioagents in the database. Identification of one or more genotypes by the methods provided herein identifies a population genotype for a population of bioagents.

[0018] In one embodiment, the population of bioagents comprises at least two bacteria. In a preferred embodiment, the population of bioagents comprises at least two bacteria belonging to the Staphylococcus genus. More preferably, the population comprises at least two bacteria belonging to the Staphylococcus aureus species. In one preferred aspect, at least one of the at least two bacteria is resistant to quinolone antimicrobial therapy. In another preferred aspect, at least one of the at least two bacteria is sensitive to quinolone antimicrobial therapy. In another preferred aspect, at least one of the at least two bacteria is resistant to quinolone antimicrobial therapy and at least one of the at least two bacteria is sensitive to quinolone antimicrobial therapy.

[0019] In one embodiment, an antibiotic regimen is developed that is tailored to treat the identified population genotype for the population of bioagents. In a preferred aspect, the antibiotic

regimen tailored to treat the identified genotypes for the population of bioagents is delivered to the sample source. In a preferred embodiment, the sample source is a human subject from whom the sample was taken.

[0020] In one embodiment, the steps of the method are periodically repeated. In one aspect, the tailored antibiotic regimen is delivered continuously during the periodic repeating of the steps. In one aspect, the antibiotic regimen is modified after one or more of the periodic repeats of the steps.

[0021] Also provided, in one embodiment, are methods for reducing a population of bacteria in a person needing such a treatment. In this embodiment, the sample is obtained from a person suspected of comprising a population of bioagents. In the identifying step of this embodiment, a population genotype is identified in the person. In one aspect, the population of bioagents in the person comprises a single genotype. In another aspect, it comprises a mixed population of bioagents, comprising at least two distinct genotypes. In this embodiment, the method further comprises administering to the person an antibiotic regimen tailored to treat the identified genotypes for the population of bioagents. In this embodiment, preferably, the population of bioagents comprises a population of bacterial bioagents. In one aspect, the steps of obtaining a sample, amplifying, determining, calculating, and identifying are repeated. In one aspect, the tailored antibiotic regimen is delivered continuously during the periodic repeating of the steps. In one aspect, during one or more of the periodic repeats of the method, an emerging genotype is identified in said sample. In this aspect, preferably, the method further comprises modifying the antibiotic regimen to treat the emerging genotype. In one embodiment, the antibiotic regimen comprises an antibiotic for treating quinolone resistant bacteria. In another embodiment, the antibiotic regimen comprises an antibiotic for treating quinolone sensitive bacteria. In another embodiment, the antibiotic regimen comprises an antibiotic for treating quinolone resistant bacteria and an antibiotic for treating quinolone sensitive bacteria. In one aspect, the antibiotic for treating quinolone sensitive bacteria is a quinolone. In one aspect, it is a fluoroquinolone.

[0022] Identification of a mixed population of bioagents allows for proper subsequent steps being performed on the sample. In one embodiment, the mixed population of bioagents comprises at least

two populations of bioagents; one population that is sensitive to a first antibiotic and another population that is resistant to said first antibiotic. Subsequent steps with such a population can include treatment with a combination of said first antibiotic to reduce the population of the bioagent sensitive thereto, and treatment with a second antibiotic to reduce the population of bioagent that is resistant to said first antibiotic.

In a further embodiment, comparison of experimental data from the sample with the database identifies only a single genotype for the population of bioagents in the sample. In one aspect of this embodiment, subsequent steps can include treatment of the population with a first antibiotic to which the population of bioagents with the one genotype is sensitive. Periodic processing of the sample is then performed as described above, thereby monitoring for the emergence of a population in the sample with a genotype that confers resistance to the administered first antibiotic. In a preferred embodiment, identification of such an emerging drug resistant bioagent or population of drug resistant bioagents is followed by alteration or modification of the treatment regimen to comprise either a second antibiotic or a combination of the first and the second antibiotics. Rapid identification of a population of bioagents in a sample allows for antibiotic regimens to be closely tailored for treatment of the specific bioagents in said sample. Further, the methods provided herein are able to identify bioagents or populations of bioagents that represent small percentages of the total population of bioagents in a sample. Genotypes in mixed populations can be identified with high sensitivity by PCR-ESI/MS because amplified bioagent nucleic acids having different base compositions appear in different positions in the mass spectrum. The dynamic range for mixed PCR-ESI/MS detections has previously been determined to be approximately 100:1(Hofstadler, S. A. et al., Inter. J. Mass Spectrom. (2005) 242, 23), which allows for detection of genotype variants with as low as 1% abundance in a mixed population. This ability allows early detection of emerging genotypes and emerging populations, including genotypes that confer drug resistance and drug resistant populations.

[0024] In one embodiment, one or more of the bioagents comprised in the population of bioagents represents less than 50% of the population of bioagents. In another embodiment, the one or more of the bioagents comprised in the population of bioagents represents less than 25% of the population of

bioagents. In another embodiment, one or more of the bioagents represents less than 10% of the population of bioagents. In another embodiment, one or more of the bioagents represents less than 5% of the population of bioagents. In another embodiment, one or more of the bioagents represents less than 4% of the population of bioagents. In another embodiment, one or more of the bioagents represents less than 3% of the population of bioagents. In another embodiment, one or more of the bioagents represents less than 2% of the population of bioagents. In another embodiment, one or more of the bioagents represents less than 2% of the population of bioagents represents less than 2% of the population of bioagents represents between about 1% and about 2% of the population of bioagents. In another embodiment, one or more of the bioagents represents about 1% of the population of bioagents.

[0025] In one embodiment, one or more of the genotypes identified by the method represents less than 50% of the population of bioagents. In another embodiment, one or more of the genotypes identified by the methods represents less than 25% of the population of bioagents. In another embodiment, one or more of the genotypes identified by the methods represents less than 15% of the population of bioagents. In another embodiment, one or more of the genotypes identified by the methods represents less than 10% of the population of bioagents. In another embodiment, one or more of the genotypes identified by the methods represents less than 5% of the population of bioagents. In another embodiment, one or more of the genotypes identified by the methods represents less than 4% of the population of bioagents. In another embodiment, one or more of the genotypes identified by the methods represents less than 3% of the population of bioagents. In another embodiment, one or more of the genotypes identified by the methods represents less than 2% of the population of bioagents. In another embodiment, one or more of the genotypes identified by the methods represents less than 2% of the population of bioagents. In another embodiment, one or more of the genotypes identified by the methods represents less than 2% of the population of bioagents. In another embodiment, one or more of the genotypes identified by the methods represents less than 2% of the population of bioagents. In another embodiment, one or more of the genotypes identified by the methods represents less than 2% of the population of bioagents. In another embodiment, one or more of the genotypes identified by the methods represents less than 2% of the population of bioagents.

BRIEF DESCRIPTION OF THE DRAWINGS

[0026] The foregoing summary and detailed description is better understood when read in conjunction with the accompanying drawings which are included by way of example and not by way of limitation.

100271 Figure 1 is a process diagram illustrating a representative primer selection process.

[0028] Figure 2 is a chart showing distribution of Staphylococcus aureus strain identification for 362 clinical isolates obtained using the genotyping primer pair panel and methods described in Example 9.

[0029] Figure 3 shows three spectra obtained using the gyrA primer pair described in Example 13. The top spectrum was generated from a patient (wound) sample, and the bottom two spectra were generated from two different colonies grown from the patient sample. In all spectra, the left peak (or double peak) represents the forward strand of the amplicon, while the right peak (or double peak) represents the reverse strand. The double peaks in the top spectrum are indicative of two different gyrA genotypes present in the patient sample. Thus, the patient sample comprised a mixed population of bioagents. As indicated by dotted lines, one peak in each of the double-peaks corresponds with the middle spectrum, representing a quinolone resistant genotype (Quinolone resistant colony gyrA mutant Ser84-Leu TCA (S) --> TTA (L)), while the other corresponds with the bottom spectrum, representing a quinolone sensitive genotype (Quinolone sensitive colony gyrA wild-type Ser84 TCA). The identification of both quinolone resistant (middle spectrum) and sensitive (bottom spectrum) genotype colonies grown from the sample is further evidence that the double peaks in the top spectrum represent a mixed population in the patient sample. Base compositions determined in this example for each amplicon are shown above each spectrum.

DETAILED DESCRIPTION OF EMBODIMENTS

[0031] As is used herein, a "bioagent" refers to any microorganism or infectious substance, or any naturally occurring, bioengineered or synthesized component of any such microorganism or infectious substance or any nucleic acid derived from any such microorganism or infectious substance. Those of ordinary skill in the art will understand fully what is meant by the term bioagent given the instant disclosure. Preferably, the bioagent is a bacterial bioagent, a bacterium or a nucleic acid derived therefrom. More preferably, the bioagent is a member of the Stophylococcus.

genus. More preferably still the bioagent is a strain of Staphylococcus aureus. A "population of bioagents" refers to a plurality of bioagents, or at least two bioagents. In some aspects, the population of bioagents is a "mixed population of bioagents," which comprises two or more distinguishable genotypes for a particular gene, locus or nucleotide position. In other aspects, each bioagent in the plurality of bioagents comprises a single genotype for the gene, locus, or nucleotide position.

[0032] As used herein, "primer pairs," or "oligonucleotide primer pairs" are synonymous terms referring to pairs of oligonucleotides (herein called "primers" or "oligonucleotide primers") that are configured to bind to conserved sequence regions of a bioagent nucleic acid (that is conserved among two or more bioagents) and to generate bioagent identifying amplicons. The bound primers flank an intervening variable region of the bioagent between the conserved sequence sequences. Upon amplification, the primer pairs yield amplicons that provide base composition variability between two or more bioagents. The variability of the base compositions allows for the identification of one or more individual bioagents from two or more bioagents based on the base composition distinctions. The primer pairs are also configured to generate amplicons that are amenable to molecular mass analysis. Each primer pair comprises two primer pair members. The primer pair members are a "forward primer" ("forward primer pair member," or "reverse member"). which comprises at least a percentage of sequence identity with the top strand of the reference sequence used in configuring the primer pair, and a "reverse primer" ("reverse primer pair member" or "reverse member"), which comprises at least a percentage of reverse complementarity with the top strand of the reference sequence used in configuring the primer pair. Primer pair configuration is well-known and is described in detail herein.

[0033] Primer pair nomenclature, as used herein, includes the identification of a reference sequence. For example, the forward primer for primer pair number 2740 is named GYRA_NC002953_7005-9668_221-249_F. This forward primer name indicates that the forward primer ("_F") hybridizes to residues 234-261 ("234_261") of a reference sequence, which in this case is represented by a sequence extraction of coordinates 7005-9668 (SEQ ID NO.: 8) from GenBank gi number 49484912 (corresponding to the version of genbank number NC 002953, as is

indicated by the prefix "GYRA NC002953" and cross-reference in Table 2). In the case of this primer, the reference sequence is the gene within a Staphylococcus aureus genome encoding for GyrA. Primer pair name codes for the primers provided herein are defined in Table 2, which lists gene abbreviations and GenBank gi numbers that correspond with each primer name code. Sequences of the primers are also provided. One of skill in the art will understand how to determine exact hybridization coordinates of primers with respect to GenBank sequences, given the information provided herein. The primer pairs are selected and configured; however, to hybridize with two or more bioagents. So, the reference sequence in the primer name is used merely to provide a reference, and not to indicate that the primers are selected and configured to hybridize with and generate a bioagent identifying amplicon only from the reference sequence. Rather, the primers hybridize with and generate amplicons from a number of sequences. Further, the sequences of the primer members of the primer pairs are not necessarily fully complementary to the conserved region of the reference bioagent. Rather, the sequences are configured to be "best fit" amongst a plurality of bioagents at these conserved binding sequences. Therefore, the primer members of the primer pairs have substantial complementarity with the conserved regions of the bioagents. including the reference bioagent.

[0034] Methods for PCR primer design are well known. One of skill in the art will understand that primer pairs configured to prime amplification of a double stranded sequence are configured and named using one strand of the double stranded sequence as a reference. The forward primer is the primer of the pair that comprises full or partial sequence identity to the one strand of the sequence being used as a reference. The reverse primer is the primer of the pair that comprises reverse complementarity to the one strand of the sequence being used as a reference.

[0035] In one embodiment, the "plus" or "top" strand (the primary sequence as submitted to GenBank) of the nucleic acid to which the primers hybridize is used as a reference when designing primer pairs. In this case, the forward primer will comprise identity and the reverse primer will comprise reverse complementarity, to the sequence listed in GenBank for the reference sequence. The ordinarily skilled artisan will understand how to configure primer pairs based upon this disclosure. In some embodiments, the primer pair is configured using the "minus" or "bottom"

strand (reverse complement of the primary sequence as submitted to and listed in GenBank). In this case, the forward primer comprises sequence identity to the minus strand, and thus comprises reverse complementarity to the top strand, the sequence listed in GenBank. Similarly, in this case, the reverse primer comprises reverse complementarity to the minus strang, and thus comprises identity to the top strand.

[0036] In a preferred embodiment, the primer pairs are configured to generate an amplicon from "within a region of SEQ ID NO.: 10," which is a specific region of Genbank gi No.: 49484912, a Staphylococcus aureus nucleic acid sequence. Configuring a primer pair to generate an amplicon from "within a region" of a particular nucleic acid reference sequence means that each primer of the pair hybridizes to a portion of the reference sequence that is within that region. One of ordinary skill in the art understands that shifting the coordinates of this region within which the primers hybridize slightly, in one direction or the other, will often result in an equally effective primer pair. Armed with the instant disclosure, one of skill in the art will be able to configure such primer pairs. Thus, in the above mentioned example, a primer pair that hybridizes to a portion of Genbank gi No.: 49484912 that is within a region slightly shifted with respect to SEQ ID NO.: 10 is encompassed by this description.

[0037] As is used herein, the term "substantial complementarity" means that a primer member of a primer pair comprises between about 70%-100%, or between about 80-100%, or between about 90-100%, or between about 99-100% sequence identity with the conserved binding sequence of any given bioagent. These ranges of identity are inclusive of all whole or partial numbers embraced within the recited range numbers. For example, and not limitation, 75.667%, 82%, 91.2435% and 97% sequence identity are all numbers that fall within the above recited range of 70% to 100%, therefore forming a part of this description

[0038] As used herein, "broad range survey primers" are intelligent primers configured to identify an unknown bioagent as a member of a particular division (e.g., an order, family, class, clade, or genus). However, in some cases the broad range survey primers are also able to identify unknown bioagents at the species or sub-species level. As used herein, "division-wide primers" are intelligent

primers configured to identify a bioagent at the species level and "drill-down" primers are intelligent primers configured to identify a bioagent at the sub-species level. As used herein, the "sub-species" level of identification includes, but is not limited to, strains, subtypes, variants, and isolates. Drill-down primers are not always required for identification at the sub-species level because broad range survey intelligent primers may, in some cases provide sufficient identification resolution to accomplishing this identification objective.

[0039] As used herein, the term "conserved region" refers to the region of the bioagent nucleic acid to which the primer pair members are designed to hybridize. Preferably, the conserved region is conserved among two or more bioagents. By the term "highly conserved," it is meant that the sequence regions exhibit between about 80-100%, or between about 90-100%, or between about 95-100% identity among all, or at least 70%, at least 80%, at least 90%, at least 95%, or at least 99% of species or strains. As used herein, the term "variable region" is used to describe a region that is between the two conserved sequence regions to which the primers of a primer pair hybridize. In other words, the variable region is a region that is flanked by the bound primers of any one primer pair described herein. The region possesses distinct base compositions among at least two bioagents, such that at least one bioagent can be identified at the family, genus, species or subspecies level using the primer pairs and the methods provided herein. The degree of variability between the at least two bioagents need only be sufficient to allow for identification using mass spectrometry or base composition analysis, as described herein. Such a difference can be as slight as a single nucleotide difference occurring between two bioagents. In a preferred embodiment, the variable region is within a reference sequence that comprises an extraction sequence with coordinates 7005-9668 (SEO ID NO.: 8) of GenBank gi number: 49484912, which comprises a nucleotide sequence encoding gyrase A (GyrA). In another preferred embodiment, the variable region is within the ORDR segment of a gene encoding gyrase A in Staphlylococcus aureus. In a preferred embodiment, this ORDR segment is SEO ID NO.: 10. In another embodiment, the variable region is within a reference sequence that comprises an extraction sequence with coordinates 7032-9695 (SEO ID NO.: 9) of GenBank gi number; 57650036, which comprises a nucleotide sequence encoding gyrase A (GyrA). In another embodiment, the variable region is within a reference sequence that comprises an extraction sequence with coordinates 7005-9674

(SEQ ID NO.: 315) of GenBank gi number: 47118324, which comprises a nucleotide sequence encoding gyrase A (GyrA). In another embodiment, the variable region is within a reference sequence that comprises an extraction sequence with coordinates 6916-9597 (SEQ ID NO.: 316) of GenBank gi number: 27314460, which comprises a nucleotide sequence encoding gyrase A (GyrA). In another preferred embodiment the variable region comprises nucleotide position 251 of a gyrA gene in Staphlylococcus aureus. In one aspect, the variable region comprises nucleotide position 251 of the reference sequence that comprises a sequence extraction with coordinates 7005-9668 (SEQ ID NO.: 8) of GenBank gi number: 49484912, which comprises a nucleotide sequence encoding Staphylococcus aureus GyrA.

[0040] As used herein, the terms "amplicon" and "bioagent identifying amplicon" refer to a nucleic acid generated using the primer pairs described herein. The amplicon is preferably double stranded DNA; however, it may be RNA and/or DNA:RNA. The amplicon comprises the sequences of the conserved regions/primer pairs and the intervening variable region. Mass spectrometry analysis of the amplicon determines a molecular mass that can be converted into a base composition. or base composition signature for the amplicon. Since the primer pairs provided herein are configured such that two or more different bioagents, when amplified with a given primer pair, will yield amplicons with unique base composition signatures, the base composition signatures can be used to identify bioagents based on association with amplicons. As discussed herein, primer pairs are configured to generate amplicons from two or more bioagents. As such, the base composition of any given amplicon will include the primer pair, the complement of the primer pair, the conserved regions and the variable region from the bioagent that was amplified to generate the amplicon. One skilled in the art understands that the incorporation of the configured primer pair sequences into any amplicon will replace the native bioagent sequences at the primer binding site, and complement thereof. After amplification of the target region using the primers the resultant amplicons having the primer sequences generate the molecular mass data. Amplicons having any native bioagent sequences at the primer binding sites, or complement thereof, are undetectable because of their low abundance. Such is accounted for when identifying one or more bioagents using any particular primer pair. The amplicon further comprises a length that is compatible with mass spectrometry

analysis. In one embodiment, bioagent identifying amplicons generate base composition signatures that are unique to the identity or genotype of a bioagent.

[0041] Calculation of base composition from a mass spectrometer generated molecular mass becomes increasingly more complex as the length of the amplicon increases. For amplicons comprising unmodified nucleic acid, the upper length as a practical length limit is about 200 consecutive nucleobases. Incorporating modified nucleotides into the amplicon can allow for an increase in this upper limit. In one embodiment, the amplicons generated using any single primer pair will provide sufficient base composition information to allow for identification of at least one bioagent at the family, genus, species or subspecies level. Alternatively, amplicons greater than 200 nucleobases can be generated and then digested to form two or more fragments that are less than 200 nucleobases. Analysis of one or more of the fragments will provide sufficient base composition information to allow for identification of at least one bioagent.

[0042] Preferably, amplicons comprise from about 45 to about 200 consecutive nucleobases (i.e., from about 45 to about 200 linked nucleosides). One of ordinary skill in the art will appreciate that this range expressly embodies compounds of 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 19, 29, 39, 49, 59, 69, 79, 89, 90, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 114, 115, 116, 117, 118, 119, 120, 121, 122, 123, 124, 125, 126, 127, 128, 129, 130, 131, 132, 133, 134, 135, 136, 137, 138, 139, 140, 141, 142, 143, 144, 145, 146, 147, 148, 149, 150, 151, 152, 153, 154, 155, 156, 157, 158, 159, 160, 161, 162, 163, 164, 165, 166, 167, 168, 169, 170, 171, 172, 173, 174, 175, 176, 177, 178, 179, 180, 181, 182, 183, 184, 185, 186, 187, 188, 189, 190, 191, 192, 193, 194, 195, 196, 197, 198, 199, and 200 nucleobases in length. One ordinarily skilled in the art will further appreciate that the above range is not an absolute limit to the length of an amplicon, but instead represents a preferred length range. Amplicons lengths falling outside of this range are also included herein so long as the amplicon is amenable to calculation of a base composition signature as herein described.

[0043] As is used herein, the term "unknown bioagent" can mean either: (i) a bioagent whose existence is not known (for example, the SARS coronavirus was unknown prior to April 2003), which is also called a "true unknown bioagent," and/or (ii) a bioagent whose existence is known (such as the well known bacterial species Staphylococcus aureus for example) but which is not known to be in a sample to be analyzed and/or (iii) a bioagent that is known or suspected of being present in a sample but whose sub-species characteristics are not known (such as a bacterial resistance genotype like the ORDR region of Staphyoicoccus aureus species). For example, if the method for identification of coronaviruses disclosed in commonly owned U.S. Pre-Grant Publication No. US2005-0266397 (incorporated herein by reference in its entirety) was to be employed prior to April 2003 to identify the SARS coronavirus in a clinical sample, both meanings of "unknown" bioagent are applicable since the SARS coronavirus was unknown to science prior to April, 2003 and since it was not known what bioagent (in this case a coronavirus) was present in the sample. On the other hand, if the method of U.S. Pre-Grant Publication No. US2005-0266397 was to be employed subsequent to April 2003 to identify the SARS coronavirus in a clinical sample, only the second meaning (ii) of "unknown" bioagent would apply because the SARS coronavirus became known to science subsequent to April 2003 but because it was not known what bioagent was present in the sample.

[0044] As used herein, the term "molecular mass" refers to the mass of a compound as determined using mass spectrometry. Herein, the compound is preferably a nucleic acid, more preferably a double stranded nucleic acid, still more preferably a double stranded DNA nucleic acid and is most preferably an amplicon. When the nucleic acid is double stranded the molecular mass is determined for both strands. Here, the strands are separated either before introduction into the mass spectrometer, or the strands are separated by the mass spectrometer (for example, electro-spray ionization will separate the hybridized strands). The molecular mass of each strand is measured by the mass spectrometer.

[0045] As used herein, the term "base composition" refers to the number of each residue comprising an amplicon, without consideration for the linear arrangement of these residues in the strand(s) of the amplicon. The amplicon residues comprise, adenosine (A), guanosine (G), cytidine,

(C), (deoxy)thymidine (T), uracil (U), inosine (D), nitroindoles such as 5-nitroindole or 3nitropyrrole, dP or dK (Hill et al.), an acyclic nucleoside analog containing 5-nitroindazole (Van Aerschot et al., Nucleosides and Nucleotides, 1995, 14, 1053-1056), the purine analog 1-(2-deoxy-.beta.-D-ribofuranosyl)-imidazole-4-carboxamide, 2,6-diaminopurine, 5-propynyluracil, 5propynylcytosine, phenoxazines, including G-clamp, 5-propynyl deoxy-cytidine, deoxy-thymidine nucleotides, 5-propynylcytidine, 5-propynyluridine and mass tag modified versions thereof, including 7-deaza-2'-deoxyadenosine-5-triphosphate, 5-jodo-2'-deoxyuridine-5'-triphosphate, 5bromo-2'-deoxyuridine-5'-triphosphate, 5-bromo-2'-deoxycytidine-5'-triphosphate, 5-iodo-2'deoxycytidine-5'-triphosphate, 5-hydroxy-2'-deoxyuridine-5'-triphosphate, 4-thiothymidine-5'triphosphate, 5-aza-2'-deoxyuridine-5'-triphosphate, 5-fluoro-2'-deoxyuridine-5'-triphosphate, O6methyl-2'-deoxyguanosine-5'-triphosphate, N2-methyl-2'-deoxyguanosine-5'-triphosphate, 8-oxo-2'deoxyguanosine-5'-triphosphate or thiothymidine-5'-triphosphate. In some embodiments, the massmodified nucleobase comprises 15.sup, N or 13.sup, C or both 15.sup, N and 13.sup, C. Preferably, the non-natural nucleosides used herein include 5-propynyluracil, 5-propynylcytosine and inosine. Herein the base composition for an unmodified DNA amplicon is notated as A.sub.wG.sub.xC.sub.vT.sub.z, wherein w, x, v and z are each independently a whole number representing the number of said nucleoside residues in an amplicon. Base compositions for amplicons comprising modified nucleosides are similarly notated to indicate the number of said natural and modified nucleosides in an amplicon. Base compositions are calculated from a molecular mass measurement of an amplicon, as described below. The calculated base composition for any given amplicon is then compared to a database of base compositions. A match between the calculated base composition and a single database entry reveals the identity of the bioagent.

[0046] As is used herein, the term "base composition signature" refers to the base composition generated by any one particular amplicon. The base composition signature for each of one or more amplicons provides a fingerprint for identifying the bioagent(s) present in a sample. Base composition signatures are unique for each genotype of the bioagent.

[0047] As used herein, the term "database" is used to refer to a collection of base composition and/or molecular mass data. The base composition and/or molecular mass data in the database is

indexed to bioagents and to primer pairs. The base composition data reported in the database comprises the number of each nucleoside in an amplicon that would be generated for each bioagent using each primer pair. The database can be populated by empirical data. In this aspect of populating the database, a bioagent is selected and a primer pair is used to generate an amplicon. The amplicon's molecular mass is determined using a mass spectrometer and the base composition calculated therefrom. An entry in the database is made to associate the base composition and/or molecular mass with the bioagent and the primer pair used. The database may also be populated using other databases comprising bioagent information. For example, using the GenBank database it is possible to perform electronic PCR using an electronic representation of a primer pair. This *in silico* method will provide the base composition for any or all selected bioagent(s) stored in the GenBank database. The information is then used to populate the base composition database as described above. A base composition database can be *in silico*, a written table, a reference book, a spreadsheet or any form generally amenable to databases. Preferably, it is *in silico*. The database can similarly be populated with molecular masses that is gathered either empirically or is calculated from other sources such as GenBank.

[0048] As used herein, the term "nucleobase" is synonymous with other terms in use in the art including "nucleotide," "deoxynucleotide," "nucleotide residue," "deoxynucleotide residue," "nucleotide triphosphate (NTP)," "residue," or deoxynucleotide triphosphate (dNTP). As is used herein, a nucleobase includes natural and modified residues, as described herein.

[0049] As used herein, a "wobble base" is a variation in a codon found at the third nucleotide position of a DNA triplet. Variations in conserved regions of sequence are often found at the third nucleotide position due to redundancy in the amino acid code.

[0050] As used herein, "housekeeping gene" refers to a gene encoding a protein or RNA involved in basic functions required for survival and reproduction of a bioagent. Housekeeping genes include, but are not limited to, genes encoding RNA or proteins involved in translation, replication, recombination and repair, transcription, nucleotide metabolism, amino acid metabolism, lipid

metabolism, energy generation, uptake, secretion and the like. In some embodiments, the primers are configured to produce amplicons from within a housekeeping gene.

[0051] As used herein, a "bioagent division" is defined as group of bioagents above the species level and includes but is not limited to, orders, families, genus, classes, clades, genera or other such groupings of bioagents above the species level.

[0052] As used herein, a "sub-species characteristic" is a genetic characteristic that provides the means to distinguish two members of the same bioagent species. For example, one bacterial strain could be distinguished from another bacterial strain of the same species by possessing a genetic change (e.g., for example, a nucleotide deletion, addition or substitution) in one of the bacterial genes, such as the GyrA gene.

[0053] As used herein, "triangulation identification" means the employment of more than one primer pair to generate a corresponding amplicon for identification of a bioagent. The more than one primer pair can be used in individual wells or in a multiplex PCR assay. Alternatively, PCR reaction may be carried out in single wells comprising a different primer pair in each well. Following amplification, the amplicons are pooled into a single well or container which is then subjected to molecular mass analysis. The combination of pooled amplicons can be chosen such that the expected ranges of molecular masses of individual amplicons are not overlapping and thus will not complicate identification of signals. Triangulation works as a process of elimination, wherein a first primer pair identifies that an unknown bioagent may be one of a group of bioagents. Subsequent primer pairs are used in triangulation identification to further refine the identity of the bioagent amongst the subset of possibilities generated with the earlier primer pair. Triangulation identification is complete when the identity of the bioagent is determined. The triangulation identification process is also used to reduce false negative and false positive signals, and enable reconstruction of the origin of hybrid or otherwise engineered bioagents. For example, identification of the three part toxin genes typical of B. anthracis (Bowen et al., J. Appl. Microbiol., 1999, 87, 270-278) in the absence of the expected signatures from the B. anthracis genome would suggest a genetic engineering event.

[0054] As is used herein, the term "single primer pair identification" means that one or more bioagents can be identified using a single primer pair. A base composition signature for an amplicon may singly identify one or more bioagents.

[0055] As used herein, the term "etiology" refers to the causes or origins, of diseases or abnormal physiological conditions.

[0056] As used herein, "population genotype" refers to the one or more genotypes for a particular gene, locus, or nucleotide position that are present in a population of bioagents. In some embodiments, the population comprises a plurality of bioagents, all with a single genotype for a particular gene, locus or nucleotide position. In these embodiments, the population genotype comprises one genotype for that gene locus or position. In other embodiments, the population of bioagents is a "mixed population," in which the plurality of bioagents has at least two distinct genotypes for a particular gene, locus or nucleotide position. In this embodiment, the population genotype comprises at least two distinct genotypes for that gene, locus or position.

[0057] The term "sample" in the present specification and claims is used in its broadest sense. On the one hand it is meant to include a specimen or culture (e.g., microbiological cultures). Preferably, the sample is from a human patient suspected of having a bacterial infection, for example, a blood, tissue, or wound sample. More preferably it is a blood, tissue, or wound swab. On the other hand, it is meant to include both biological and environmental samples. A sample may include a specimen of synthetic origin. Biological samples may be from an animal, including human, and may be fluid, solid (e.g., stool) or tissue, as well as liquid or solid food and feed products or ingredients such as dairy items, vegetables, meat and meat by-products, or waste. Biological samples may be obtained from all of the various families of domestic animals, as well as feral or wild animals, including, but not limited to, such animals as ungulates, bear, fish, lagamorphs, rodents, etc. Environmental samples include environmental material such as surface matter, soil, water, air and industrial samples include environmental material such as surface matter, soil, water, air and industrial samples, as well as samples obtained from food and dairy processing instruments, apparatus, equipment, utensils, disposable and non-disposable items. These examples are not to be construed as limiting the sample types applicable to the present invention. The term "source of target nucleic

acid" refers to any sample that contains nucleic acids (RNA or DNA). Particularly preferred sources of target nucleic acids are biological samples including, but not limited to blood, saliva, cerebral spinal fluid, pleural fluid, milk, lymph, sputum and semen. In some embodiments, the sample is purified. The term "sample source" refers to the source of the sample, for example, the animal, human, fluid, tissue, culture, or other source from which the sample was isolated and/or purified.

[0058] Provided herein are methods for detection and identification of bioagents in an unbiased manner using bioagent identifying amplicons. In one aspect, the methods are for detection and identification of population genotype for a population of bioagents. Primers are selected to hybridize to conserved sequence regions of nucleic acids derived from a bioagent and which bracket (flank) variable sequence regions to yield a bioagent identifying amplicon which can be amplified and which is amenable to molecular mass determination. The molecular mass is converted to a base composition, which indicates the number of each nucleotide in the amplicon. The molecular mass or corresponding base composition signature of the amplicon is then queried against a database of molecular masses or base composition signatures indexed to bioagents and to the primer pair used to generate the amplicon. A match of the measured base composition to a database entry base composition associates the sample bioagent to an indexed bioagent in the database. Thus, the identity of the unknown bioagent or population of bioagents is determined. Prior knowledge of the unknown bioagent or population of bioagents is not necessary. In some instances, the measured base composition associates with more than one database entry base composition. Thus, a second/subsequent primer pair is used to generate an amplicon, and its measured base composition is similarly compared to the database to determine its identity in triangulation identification. For example, a first primer pair might identify that a bacterial bioagent is present in a sample that is a member of the Staphylococcus genus. A second primer might determine that it is a member of the Staphylococcus aureus species. A third primer pair might identify that the bioagent is resistant to quinolones. Furthermore, the method can be applied to rapid parallel multiplex analyses, the results of which can be employed in a triangulation identification strategy. The present method provides rapid throughput and does not require nucleic acid sequencing of the amplified target sequence for bioagent detection and identification.

[0059] In some embodiments, the methods are performed on nucleic acids comprised in a sample suspected of comprising a population of bioagents. In one aspect, the methods further comprise administering or delivering to the sample source an antibiotic regimen tailored to treat the identified genotypes for the population of bacteria. In this aspect, the antibiotic regimen is determined based on the genotype(s) identified by the method, with the goal of being able to effectively reduce the bioagents in the population. In one embodiment, the steps of the method are repeated "periodically" or more than one additional time following the initial identification. In one aspect, the periodic repeating of the steps is done at regular intervals. In other aspects, it is done sporadically or at irregular time points. In another aspect, it is done in response to a trigger, such as the appearance of one or more symptoms. In one aspect, the antibiotic regimen is modified based on one or more genotypes identified during the periodic repeating of the steps. In one embodiment, the antibiotic regimen comprises an antibiotic for treating quinolone resistant bacteria. In another embodiment, the antibiotic regimen comprises an antibiotic for treating quinolone sensitive bacteria. In one aspect, it is a fluoroouinolone.

[0060] Despite enormous biological diversity, all forms of life on earth share sets of essential, common features in their genomes. Since genetic data provide the underlying basis for identification of bioagents by the current methods, it is necessary to select segments of nucleic acids which ideally provide enough variability to distinguish each individual bioagent and whose molecular mass is amenable to molecular mass determination.

[0061] In some embodiments, at least one bacterial nucleic acid segment is amplified in the process of identifying the bioagent. Thus, the nucleic acid segments that can be amplified by the primers disclosed herein and that provide enough variability to distinguish each individual bioagent and whose molecular masses are amenable to molecular mass determination are herein described as bioagent identifying amplicons.

[0062] In some embodiments, bioagent identifying amplicons amenable to molecular mass determination that are produced by the primers described herein are either of a length, size and/or

mass compatible with the particular mode of molecular mass determination or compatible with a means of providing a predictable fragmentation pattern in order to obtain predictable fragments of a length compatible with the particular mode of molecular mass determination. Such means of providing a predictable fragmentation pattern of an amplicon include, but are not limited to, cleavage with restriction enzymes or cleavage primers, for example. Thus, in some embodiments, bioagent identifying amplicons are larger than 200 nucleobases and are amenable to molecular mass determination following restriction digestion. Methods of using restriction enzymes and cleavage primers are well known to those with ordinary skill in the art.

[0063] In some embodiments, amplicons corresponding to bioagent identifying amplicons are obtained using the polymerase chain reaction (PCR) which is a routine method to those with ordinary skill in the molecular biology arts. Other amplification methods may be used such as ligase chain reaction (LCR), low-stringency single primer PCR, and multiple strand displacement amplification (MDA). These methods are also known to those with ordinary skill. (Michael, SF., Biotechniques (1994), 16:411-412 and Dean et al., Proc. Natl. Acad. Sci. U.S.A. (2002), 99, 5261-5266)

[0064] A representative process flow diagram used for primer selection and validation process is outlined in Figure 1. For each group of diverse organisms, candidate target sequences are identified (200) from which nucleotide alignments are created (210) and analyzed (220). Primers are then configured by selecting appropriate priming regions (230) to facilitate the selection of candidate primer pairs (240). The primer pair sequence is a "best fit" amongst the aligned sequences, meaning that the primer pair sequence may or may not be fully complementary to the hybridization region on any one of the bioagents in the alignment. Thus, bets fit primer pair sequences are those with sufficient complementarity with two or more bioagents to hybridize with the two or more bioagents and generate an amplicon. The primer pairs are then subjected to in silico analysis by electronic PCR (ePCR) (300) wherein bioagent identifying amplicons are obtained from sequence databases such as GenBank or other sequence collections (310) and checked for specificity in silico (320). Bioagent identifying amplicons obtained from ePCR of GenBank sequences (310) can also be analyzed by a probability model which predicts the capability of a given amplicon to identify

unknown bioagents. Preferably, the base compositions of amplicons with favorable probability scores are then stored in a base composition database (325). Alternatively, base compositions of the bioagent identifying amplicons obtained from the primers and GenBank sequences can be directly entered into the base composition database (330). Candidate primer pairs (240) are validated by in vitro amplification by a method such as PCR analysis (400) of nucleic acid from a collection of organisms (410). Amplicons thus obtained are analyzed to confirm the sensitivity, specificity and reproducibility of the primers used to obtain the amplicons (420).

[0065] Synthesis of primers is well known and routine in the art. The primers may be conveniently and routinely made through the well-known technique of solid phase synthesis. Equipment for such synthesis is sold by several vendors including, for example, Applied Biosystems (Foster City, CA). Any other means for such synthesis known in the art may additionally or alternatively be employed.

The primers are employed as compositions for use in methods for identification of bacterial bioagents as follows: a primer pair composition is contacted with nucleic acid (such as, for example, DNA or DNA reverse transcribed from RNA) of an unknown bacterial bioagent. The nucleic acid is then amplified by a nucleic acid amplification technique, such as PCR for example, to obtain an amplicon that represents a bioagent identifying amplicon. The molecular mass of each strand of the double-stranded amplicon is determined by a molecular mass measurement technique such as mass spectrometry for example. Preferably the two strands of the double-stranded amplicon are separated during the ionization process; however, they may be separated prior to mass spectrometry measurement. In some embodiments, the mass spectrometer is electrospray Fourier transform ion cyclotron resonance mass spectrometry (ESI-FTICR-MS) or electrospray time of flight mass spectrometry (ESI-TOF-MS). A list of possible base compositions can be generated for the molecular mass value obtained for each strand and the choice of the correct base composition from the list is facilitated by matching the base composition of one strand with a complementary base composition of the other strand. The measured molecular mass or base composition calculated therefrom is then compared with or querried against a database of molecular masses or base compositions indexed to primer pairs and to known bacterial bioagents. A match between the

measured molecular mass or base composition of the amplicon and the database molecular mass or base composition for that indexed primer pair will associate the measured molecular mass or base composition with an indexed bacterial bioagent, thus indicating the identity of the unknown bioagent. In some embodiments, the primer pair used is one of the primer pairs of Table 1. In some embodiments, the method is repeated using a different primer pair to resolve possible ambiguities in the identification process or to improve the confidence level for the identification assignment (triangulation identification).

[0067] In some embodiments, a bioagent identifying amplicon may be produced using only a single primer (either the forward or reverse primer of any given primer pair), provided an appropriate amplification method is chosen, such as, for example, low stringency single primer PCR (LSSP-PCR). Adaptation of this amplification method in order to produce bioagent identifying amplicons can be accomplished by one with ordinary skill in the art without undue experimentation. (Pena, SDJ et al., Proc. Natl. Acad. Sci. U.S.A (1994) 91, 1946–1949).

[0068] In some embodiments, the oligonucleotide primers are broad range survey primers which hybridize to conserved regions of a nucleic acid encoding a gene that is common to all known members of the Staphylococcus genus, though the sequences of the gene that are within the variable region vary. The broad range primer may identify the unknown bioagent, depending on which bioagent is in the sample. In other cases, the molecular mass or base composition of an amplicon does not provide enough resolution to unambiguously identify the unknown bioagent as any one bacterial bioagent at or below the species level. These cases benefit from further analysis of one or more an amplicons generated from at least one additional broad range survey primer pair or from at least one additional division-wide primer pair or from at least one additional division of sub-species characteristics is often critical for determining proper clinical treatment of viral infections, or in rapidly responding to an outbreak of a new viral strain to prevent massive epidemic or pandemic.

[0069] In some embodiments, the primers used for amplification hybridize to and amplify genomic DNA, DNA of bacterial plasmids, transposons and other exogenous nucleic acid, or DNA

reverse transcribed from RNA. Among other things, the identification of non-bacterial nucleic acids or combinations of bacterial and non-bacterial nucleic acids are useful for detecting bioengineered bioagents.

[0070] In some embodiments, the primers used for amplification hybridize directly to bacterial RNA and act as reverse transcription primers for obtaining DNA from direct amplification of bacterial RNA. Methods of amplifying RNA to produce cDNA using reverse transcriptase are well known to those with ordinary skill in the art and can be routinely established without undue experimentation.

[0071] One with ordinary skill in the art of design of amplification primers will recognize that a given primer need not hybridize with 100% complementarity in order to effectively prime the synthesis of a complementary nucleic acid strand in an amplification reaction. Primer pair sequences may be a "best fit" amongst the aligned bioagent sequences, thus not be fully complementary to the hybridization region on any one of the bioagents in the alignment. Moreover, a primer may hybridize over one or more segments such that intervening or adjacent segments are not involved in the hybridization event. (e.g., for example, a loop structure or a hairpin structure). The primers may comprise at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95% or at least 99% sequence identity with any of the primers listed in Table 1 or other primer disclosed herein. Thus, in some embodiments, an extent of variation of 70% to 100%, or any range falling within, of the sequence identity is possible relative to the specific primer sequences disclosed herein. Determination of sequence identity is described in the following example: a primer 20 nucleobases in length which is identical to another 20 nucleobase primer having two non-identical residues has 18 of 20 identical residues (18/20 = 0.9 or 90% sequence identity). In another example, a primer 15 nucleobases in length having all residues identical to a 15 nucleobase segment of primer 20 nucleobases in length would have 15/20 = 0.75 or 75% sequence identity with the 20 nucleobase primer. Percent identity need not be a whole number, for example when a 28 consecutive nucleobase primer is completely identical to a 31 consecutive nucleobase primer (28/31 = 0.9032 or 90.3% identical). Similarly, either or both of the primers of the primer pairs provided herein may comprise 0-9 nucleobase deletions, additions, and/or substitutions relative to any of the primers

listed in Table 1, or elsewhere herein. In other words, either or both of the primers may comprise 0, 1, 2, 3, 4, 5, 6, 7, 8 or 9 nucleobase deletions, 0, 1, 2, 3, 4, 5, 6, 7, 8 or 9 nucleobase additions, 0, 1, 2, 3, 4, 5, 6, 7, 8 or 9 nucleobase substitutions relative to the sequences of any of the primers disclosed herein. In one aspect, the primers comprise the sequence of any of the primers listed in Table 1 with the non-templated T residue removed from the 5' terminus. In one aspect, the primers comprise the sequence of any of the primers listed in Table 1 with the non-templated T residue removed from the 5' terminus and comprising 0-9 nucleobase deletions, additions, and/or substitutions.

[0072] Percent homology, sequence identity or target complementarity, can be determined by, for example, the Gap program (Wisconsin Sequence Analysis Package, Version 8 for Unix, Genetics Computer Group, University Research Park, Madison WI), using default settings, which uses the algorithm of Smith and Waterman (Adv. Appl. Math., 1981, 2, 482-489). In some embodiments, target complementarity of primers with respect to the conserved priming regions of bacterial nucleic acid, is between about 70% and about 80%. In other embodiments, homology, sequence identity or complementarity, is between about 80% and about 90%. In yet other embodiments, homology, sequence identity or complementarity, is at least 90%, at least 92%, at least 94%, at least 95%, at least 95%, at least 95% at least 95% at least 95% at least 95%.

[0073] In some embodiments, the primers described herein comprise at least 70%, at least 75%, at least 80%, at least 80%, at least 90%, at least 92%, at least 94%, at least 95%, at least 98%, or at least 99%, or 100% (or any range falling within) sequence identity with the primer sequences specifically disclosed herein.

[0074] One with ordinary skill is able to calculate percent sequence identity or percent sequence homology and is able to determine, without undue experimentation, the effects of variation of primer sequence identity on the function of the primer in its role in priming synthesis of a complementary strand of nucleic acid for production of a corresponding bioagent identifying amplicon.

[0075] In some embodiments, the oligonucleotide primers are 13 to 35 nucleobases in length (13 to 35 linked nucleotide residues). These embodiments comprise oligonucleotide primers 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34 or 35 nucleobases in length, or any range therewithin.

[0076] In some embodiments, any given primer comprises a modification comprising the addition of a non-templated T residue to the 5' end of the primer (i.e., the added T residue does not necessarily hybridize to the nucleic acid being amplified). The addition of a non-templated T residue has an effect of minimizing the addition of non-templated A residues as a result of the non-specific enzyme activity of *Taq* polymerase (Magnuson *et al.*, Biotechniques, 1996, 21, 700-709), an occurrence which may lead to ambiguous results arising from molecular mass analysis. Primer pairs comprising the sequence of any of the primer pairs described herein, but lacking the non-templated T residue at the 5' end of the primer are also encompassed by this disclosure.

[0077] Primers may contain one or more universal bases. Because any variation (due to codon wobble in the third position) in the conserved regions among species is likely to occur in the third position of a DNA (or RNA) triplet, oligonucleotide primers can be configured such that the nucleotide corresponding to this position is a base which can bind to more than one nucleotide, referred to herein as a "universal nucleobase." For example, under this "wobble" pairing, inosine (I) binds to U, C or A; guanine (G) binds to U or C, and uridine (U) binds to U or C. Other examples of universal nucleobases include nitroindoles such as 5-nitroindole or 3-nitropytrole (Loakes et al., Nucleosides and Nucleotides, 1995, 14, 1001-1003), the degenerate nucleotides dP or dK (Hill et al.), an acyclic nucleoside analog containing 5-nitroindazole (Van Aerschot et al., Nucleosides and Nucleotides, 1995, 14, 1033-1056) or the purine analog 1-(2-deoxy-beta. -D-ribofuranosyl)-imidazole-4-carboxamide (Sala et al., Nucl. Acids Res., 1996, 24, 3302-3306)

[0078] In some embodiments, to compensate for the somewhat weaker binding by the wobble base, the oligonucleotide primers are configured such that the first and second positions of each triplet are occupied by nucleotide analogs which bind with greater affinity than the unmodified nucleotide. Examples of these analogs include, but are not limited to, 2,6-diaminopurine which

binds to thymine, 5-propynyluracil which binds to adenine and 5-propynylcytosine and phenoxazines, including G-clamp, which binds to G. Propynylated pyrimidines are described in U.S. Patent Nos. 5,645,985, 5,830,653 and 5,484,908, each of which is commonly owned and incorporated herein by reference in its entirety. Propynylated primers are described in U.S. Pre-Grant Publication No. 2003-0170682; also commonly owned and incorporated herein by reference in its entirety. Phenoxazines are described in U.S. Patent Nos. 5,502,177, 5,763,588, and 6,005,096, each of which is incorporated herein by reference in its entirety. G-clamps are described in U.S. Patent Nos. 6,007,992 and 6,028,183, each of which is incorporated herein by reference in its entirety.

[0079] In some embodiments, to enable broad priming of rapidly evolving bioagents, primer hybridization is enhanced using primers and probes containing 5-propynyl deoxy-cytidine and deoxy-thymidine nucleotides. These modified primers offer increased affinity and base pairing selectivity.

[0080] In some embodiments, non-template primer tags are used to increase the melting temperature (T.sub.m) of a primer-template duplex in order to improve amplification efficiency. A non-template tag is at least three consecutive A or T nucleotide residues on a primer which are not complementary to the template. In any given non-template tag, A can be replaced by C or G and T can also be replaced by C or G. Although Watson-Crick hybridization is not expected to occur for a non-template tag relative to the template, the extra hydrogen bond in a G-C pair relative to an A-T pair confers increased stability of the primer-template duplex and improves amplification efficiency for subsequent cycles of amplification when the primers hybridize to strands synthesized in previous cycles.

[0081] In other embodiments, propynylated tags may be used in a manner similar to that of the non-template tag, wherein two or more 5-propynylcytidine or 5-propynyluridine residues replace template matching residues on a primer. In other embodiments, a primer contains a modified internucleoside linkage such as a phosphorothioate linkage, for example.

[0082] In some embodiments, the primers contain mass-modifying tags. Reducing the total number of possible base compositions of a nucleic acid of specific molecular weight provides a means of avoiding a persistent source of ambiguity in determination of base composition of amplicons. Addition of mass-modifying tags to certain nucleobases of a given primer will result in simplification of *de novo* determination of base composition of a given bioagent identifying amplicon from its molecular mass.

[0083] In some embodiments, the mass modified nucleobase comprises one or more of the following: for example, 7-deaza-2'-deoxyadenosine-5-triphosphate, 5-iodo-2'-deoxyuridine-5'-triphosphate, 5-bromo-2'-deoxycytidine-5'-triphosphate, 5-iodo-2'-deoxycytidine-5'-triphosphate, 5-iodo-2'-deoxycytidine-5'-triphosphate, 5-iodo-2'-deoxycytidine-5'-triphosphate, 5-hydroxy-2'-deoxyuridine-5'-triphosphate, 4-thiothymidine-5'-triphosphate, 5-aza-2'-deoxyuridine-5'-triphosphate, 5-fluoro-2'-deoxyuridine-5'-triphosphate, N2-methyl-2'-deoxyguanosine-5'-triphosphate, N2-methyl-2'-deoxyguanosine-5'-triphosphate or thiothymidine-5'-triphosphate. In some embodiments, the mass-modified nucleobase comprises sup.15N or sup.13C or both sup.15N and sup.13C.

[0084] In some embodiments, the molecular mass of a given bioagent identifying amplicon is determined by mass spectrometry. Mass spectrometry has several advantages, not the least of which is high bandwidth characterized by the ability to separate (and isolate) many molecular peaks across a broad range of mass to charge ratio (m/z). Thus mass spectrometry is intrinsically a parallel detection scheme without the need for radioactive or fluorescent labels since every amplicon is identified by its molecular mass. The current state of the art in mass spectrometry is such that less than femtomole quantities of material can be readily analyzed to afford information about the molecular contents of the sample. An accurate assessment of the molecular mass of the material can be quickly obtained, irrespective of whether the molecular weight of the sample is several hundred, or in excess of one hundred thousand atomic mass units (amu) or Daltons.

[0085] In some embodiments, intact molecular ions are generated from amplicons using one of a variety of ionization techniques to convert the sample to gas phase. These ionization methods

include, but are not limited to, electrospray ionization (ES), matrix-assisted laser desorption ionization (MALDI) and fast atom bombardment (FAB). Upon ionization, several peaks are observed from one sample due to the formation of ions with different charges. Averaging the multiple readings of molecular mass obtained from a single mass spectrum affords an estimate of molecular mass of the bioagent identifying amplicon. Electrospray ionization mass spectrometry (ESI-MS) is particularly useful for very high molecular weight polymers such as proteins and nucleic acids having molecular weights greater than 10 kDa, since it yields a distribution of multiply-charged molecules of the sample without causing a significant amount of fragmentation.

[0086] The mass detectors used include, but are not limited to, Fourier transform ion cyclotron resonance mass spectrometry (FT-ICR-MS), time of flight (TOF), ion trap, quadrupole, magnetic sector, Q-TOF, and triple quadrupole.

[0087] In some embodiments, assignment of previously unobserved base compositions (also known as "true unknown base compositions") to a given phylogeny can be accomplished via the use of pattern classifier model algorithms. Base compositions, like sequences, vary slightly from strain to strain within species, for example. In some embodiments, the pattern classifier model is the mutational probability model. On other embodiments, the pattern classifier is the polytope model.

[0088] In one embodiment, it is possible to manage this diversity by building "base composition probability clouds" around the composition constraints for each species. This permits identification of organisms in a fashion similar to sequence analysis. Using three primer pairs, a "pseudo four-dimensional plot" can be used to visualize the concept of base composition probability clouds. Optimal primer design requires optimal choice of bioagent identifying amplicons and maximizes the separation between the base composition signatures of individual bioagents. Areas where clouds overlap indicate regions that may result in a misclassification, a problem which is overcome by a triangulation identification process using bioagent identifying amplicons not affected by overlap of base composition probability clouds.

[0089] In some embodiments, base composition probability clouds provide the means for screening potential primer pairs in order to avoid potential misclassifications of base compositions. In other embodiments, base composition probability clouds provide the means for predicting the identity of an unknown bioagent whose assigned base composition was not previously observed and/or indexed in a bioagent identifying amplicon base composition database due to evolutionary transitions in its nucleic acid sequence. Thus, in contrast to probe-based techniques, mass spectrometry determination of base composition does not require prior knowledge of the composition or sequence in order to make the measurement.

[0090] Provided herein are bioagent classifying information at a level sufficient to identify a given bioagent. Furthermore, the process of determining a previously unknown base composition for a given bioagent (for example, in a case where sequence information is unavailable) has downstream utility by providing additional bioagent indexing information with which to populate base composition databases. The process of future bioagent identification is thus greatly improved as more base composition signature indexes become available in base composition databases.

[0091] In some embodiments, the identity and quantity of an unknown bioagent can be determined using the process illustrated in Figure 4. Primers (500) and a known quantity of a calibration polynucleotide (505) is added to a sample containing nucleic acid of an unknown bioagent. The total nucleic acid in the sample is then subjected to an amplification reaction (510) to obtain amplicons. The molecular masses of amplicons are determined (515) from which are obtained molecular mass and abundance data. The molecular mass of the bioagent identifying amplicon (520) provides for its identification (525) and the molecular mass of the calibration amplicon obtained from the calibration polynucleotide (530) provides for its quantification (535). The abundance data of the bioagent identifying amplicon is recorded (540) and the abundance data for the calibration data is recorded (545), both of which are used in a calculation (550) which determines the quantity of unknown bioagent in the sample.

[0092] A sample comprising an unknown bioagent is contacted with a primer pair which amplifies the nucleic acid from the bioagent, and a known quantity of a polynucleotide that

comprises a calibration sequence. The rate of amplification is reasonably assumed to be similar for the nucleic acid of the bioagent and for the calibration sequence. The amplification reaction then produces two amplicons: a bioagent identifying amplicon and a calibration amplicon. The bioagent identifying amplicon and the calibration amplicon should be distinguishable by molecular mass while being amplified at essentially the same rate. Effecting differential molecular masses can be accomplished by choosing as a calibration sequence, a representative bioagent identifying amplicon (from a specific species of bioagent) and performing, for example, a 2-8 nucleobase deletion or insertion within the variable region between the two priming sites. The amplified sample containing the bioagent identifying amplicon and the calibration amplicon is then subjected to molecular mass analysis by mass spectrometry, for example. The resulting molecular mass analysis of the nucleic acid of the bioagent and of the calibration sequence. The molecular mass data obtained for the nucleic acid of the bioagent and of the calibration sequence. The molecular mass data obtained for the nucleic acid of the bioagent and sequence and the quantity of the bioagent, based on the knowledge of the quantity of calibration polynucleotide contacted with the sample.

[0093] In some embodiments, construction of a standard curve where the amount of calibration polynucleotide spiked into the sample is varied provides additional resolution and improved confidence for the determination of the quantity of bioagent in the sample. The use of standard curves for analytical determination of molecular quantities is well known to one with ordinary skill and can be performed without undue experimentation. Alternatively, the calibration polynucleotide can be amplified in into own reaction well or wells under the same conditions as the bioagent. A standard curve can be prepared therefrom, and a relative abundance of the bioagent determined by methods such as linear regression. In some embodiments, multiplex amplification is performed where multiple bioagent identifying amplicons are amplified with multiple primer pairs which also amplify the corresponding standard calibration sequences. In this or other embodiments, the standard calibration sequences are optionally included within a single construct (preferably a vector) which functions as the calibration polynucleotide. Competitive PCR, quantitative PCR, quantitative competitive PCR, multiplex and calibration polynucleotides are all methods and materials well known to those ordinarily skilled in the art and can be performed without undue experimentation.

[0094] In some embodiments, the calibrant polynucleotide is used as an internal positive control to confirm that amplification conditions and subsequent analysis steps are successful in producing a measurable amplicon. Even in the absence of copies of the genome of a bioagent, the calibration polynucleotide should give rise to a calibration amplicon. Failure to produce a measurable calibration amplicon indicates a failure of amplification or subsequent analysis step such as amplicon purification or molecular mass determination. Reaching a conclusion that such failures have occurred is in itself, a useful event. In some embodiments, the calibration sequence is comprised of DNA. In some embodiments, the calibration sequence is comprised of PNA.

[0095] In the preferred embodiment, the calibration sequence is inserted into a vector which then itself functions as the calibration polynucleotide. In some embodiments, more than one calibration sequence is inserted into the vector that functions as the calibration polynucleotide. Such a calibration polynucleotide is herein termed a "combination calibration polynucleotide." The process of inserting polynucleotides into vectors is routine to those skilled in the art and can be accomplished without undue experimentation. Thus, it should be recognized that the calibration method should not be limited to the embodiments described herein. The calibration method can be applied for determination of the quantity of any bioagent identifying amplicon when an appropriate standard calibrant polynucleotide sequence is configured and used. The process of choosing an appropriate vector for insertion of a calibrant is also a routine operation that can be accomplished by one with ordinary skill without undue experimentation.

[0096] It is preferable for some primer pairs to produce bioagent identifying amplicons within more conserved regions of Staphylococci bacteria while others produce bioagent identifying amplicons within regions that are likely to evolve more quickly. Primer pairs that characterize amplicons in a conserved region with low probability that the region will evolve past the point of primer recognition are useful as a broad range survey-type primer. Primer pairs that characterize an amplicon corresponding to an evolving genomic region are useful for distinguishing emerging strain variants.

[0097] The primer pairs described herein establish a platform for identifying members of the Staphylococcus genus. Base composition analysis eliminates the need for prior knowledge of bioagent sequence to generate hybridization probes. Thus, in another embodiment, there is provided a method for determining the etiology of a bacterial infection when the process of identification of bacteria is carried out in a clinical setting and, even when the bacteria is a new species never observed before. This is possible because the methods are not confounded by naturally occurring evolutionary variations (a major concern when using probe based or sequencing dependent methods for characterizing viruses that evolve rapidly). Measurement of molecular mass and determination of base composition is accomplished in an unbiased manner without sequence prejudice and without the need for specificity as is required with probes.

[0098] Another embodiment provides a means of tracking the spread of any species or strain of bacteria when a plurality of samples obtained from different locations are analyzed by the methods described above in an epidemiological setting. For example, a plurality of samples from a plurality of different locations is analyzed with primers which produce bioagent identifying amplicons, a subset of which contains a specific bacteria. The corresponding locations of the members of the bacteria-containing subset indicate the spread of the specific bacteria to the corresponding locations.

[0099] Also provided are kits for carrying out the methods described herein. In some embodiments, the kit may comprise a sufficient quantity of one or more primer pairs to perform an amplification reaction on a target polynucleotide from a bioagent to form a bioagent identifying amplicon. In some embodiments, the kit may comprise from one to fifty primer pairs, from one to twenty primer pairs, from one to ten primer pairs, from one to eight primer pairs or from two to five primer pairs. In some embodiments, the kit may comprise one or more primer pairs recited in Table 1. In a preferred embodiment, the kit comprises eight primer pairs from Table 1. In a preferred aspect the eight primer pairs comprised in the kit are selected from: SEQ ID NO: 58:SEQ ID NO: 142, SEQ ID NO: 62:SEQ ID NO: 147, SEQ ID NO: 294.SEQ ID NO: 295, SEQ ID NO: 35:SEQ ID NO: 121, SEQ ID NO: 39:SEQ ID NO: 121, SEQ ID NO: 139, SEQ ID NO: 121, SEQ ID NO: 21:SEQ ID NO: 22:SEQ ID NO: 32:SEQ ID NO: 330. SEQ ID NO: 300. SEQ ID

331:SEQ ID NO.:332, SEQ ID NO.: 2:SEQ ID NO.:5, SEQ ID NO.: 3:SEQ ID NO.:6, SEQ ID NO.: 3:SEQ ID NO.:7, and SEQ ID NO.: 4:SEQ ID NO.:5. In another preferred aspect, the eight primer pairs comprised in the kit are selected from: SEO ID NO.: 72:SEO ID NO.:156, SEO ID NO.: 79:SEO ID NO.:166. SEO ID NO.: 76:SEO ID NO.:162. SEO ID NO.: 83:SEO ID NO.:170. SEO ID NO.: 87:SEO ID NO.: 172, SEO ID NO.: 90:SEO ID NO.: 177, SEO ID NO.: 93:SEO ID NO.:180. SEO ID NO.: 94:SEO ID NO.:181. SEO ID NO.: 72:SEO ID NO.:158. SEO ID NO.: 2:SEO ID NO.:5, SEO ID NO.: 3:SEO ID NO.:6, SEO ID NO.: 3:SEO ID NO.:7, and SEO ID NO.: 4:SEO ID NO.:5. In another preferred embodiment, the kit comprises nine oligonucleotide primer pairs. In a preferred aspect, the nine oligonucleotide primer pairs are SEO ID NO.: 58:SEO ID NO.:142. SEO ID NO.: 62:SEO ID NO.:147. SEO ID NO.: 294:SEO ID NO.:295. SEO ID NO.: 35:SEO ID NO.:121, SEO ID NO.: 39:SEO ID NO.:125, SEO ID NO.: 47:SEO ID NO.:132, SEO ID NO.: 55:SEO ID NO.:139, SEO ID NO : 21:SEO ID NO.:104, SEO ID NO : 22:SEO ID NO.:106, SEO ID NO.: 70:SEO ID NO.:155, and SEO ID NO.: 3:SEO ID NO.:7. In another preferred aspect, the nine oligonucleotide primers comprised in the kit are SEO ID NO.: 72:SEO ID NO.:156, SEO ID NO.: 79:SEO ID NO.:166, SEO ID NO.: 76:SEO ID NO.:162, SEO ID NO.: 83:SEO ID NO.:170, SEO ID NO.: 87:SEO ID NO.:172, SEO ID NO.: 90:SEO ID NO.:177, SEO ID NO.: 93:SEQ ID NO.:180, SEQ ID NO.: 94:SEQ ID NO.:181, SEQ ID NO.: 72:SEQ ID NO.:158, and SEO ID NO.: 3:SEO ID NO.:7. In another preferred embodiment, the kit comprises 17 oligonucleotide primer pairs. Preferrably, the 17 oligonucleotide primer pairs comprised in the kit are SEQ ID NO.: 58:SEQ ID NO.:142, SEQ ID NO.: 62:SEQ ID NO.:147, SEQ ID NO.: 294:SEQ ID NO.:295, SEQ ID NO.: 35:SEQ ID NO.:121, SEQ ID NO.: 39:SEQ ID NO.:125, SEQ ID NO.: 47:SEQ ID NO.:132, SEQ ID NO.: 55:SEQ ID NO.:139, SEQ ID NO.: 21:SEQ ID NO.:104, SEQ ID NO.: 22:SEQ ID NO.:106, SEQ ID NO.: 70:SEQ ID NO.: 155, SEQ ID NO.: 72:SEQ ID NO.:156, SEQ ID NO.: 79:SEQ ID NO.:166, SEQ ID NO.: 76:SEQ ID NO.:162, SEQ ID NO.: 83:SEQ ID NO.:170, SEQ ID NO.: 87:SEQ ID NO.:172, SEQ ID NO.: 90:SEQ ID NO.:177, SEQ ID NO.: 93:SEQ ID NO.:180, SEQ ID NO.: 94:SEQ ID NO.:181, SEQ ID NO.: 72:SEQ ID NO.:158, and SEQ ID NO.: 3:SEQ ID NO.:7.

[00100] In some embodiments, the kit may comprise one or more broad range survey primer(s), division wide primer(s), or drill-down primer(s), or any combination thereof. A kit may be

configured so as to comprise select primer pairs for identification of a particular bioagent. For example, a broad range survey primer kit may be used initially to identify an unknown bioagent as a member of the genus Staphylococcus. Another example of a division-wide kit may be used to distinguish Staphylococcus aureus from Staphylococcus epidermidis, for example. A drill-down kit may be used, for example, to distinguish resistance and sensitivity of bacteria to one or more antibiotics. In some embodiments, the kit may contain standardized calibration polynucleotides for use as internal amplification calibrants.

[00101] In some embodiments, the kit may also comprise a sufficient quantity of reverse transcriptase (if an RNA is to be identified for example), a DNA polymerase, suitable nucleoside triphosphates (including any of those described above), a DNA ligase, and/or reaction buffer, or any combination thereof, for the amplification processes described above. A kit may further include instructions pertinent for the particular embodiment of the kit, such instructions describing the primer pairs and amplification conditions for operation of the method. A kit may also comprise amplification reaction containers such as microcentrifuge tubes and the like. A kit may also comprise reagents or other materials for isolating bioagent nucleic acid or bioagent identifying amplicons from amplification, including, for example, detergents, solvents, or ion exchange resins which may be linked to magnetic beads. A kit may also comprise a table of measured or calculated molecular masses and/or base compositions of bioagents using the primer pairs of the kit.

[00102] In one embodiment, population genotypes for mixed populations of bioagents can are identified. Population genotypes for mixed populations can be identified with high sensitivity by PCR-ESI/MS because amplified bioagent nucleic acids having different base compositions appear in different positions in the mass spectrum. The dynamic range for mixed PCR-ESI/MS detections has previously been determined to be approximately 100:1(Hofstadler, S. A. et al., Inter. J. Mass Spectrom. (2005) 242, 23), which allows for detection of genotype variants with as low as 1% abundance in a mixed population. This detection using PCR-ESI/MS surveillance does not require secondary testing.

[00103] The following examples serve only as illustration, and not limitation.

EXAMPLES

Example 1: Selection of Design and Validation of Primers that Define Bioagent Identifying Amplicons for Staphylococcus

[00104] For design of primers that define Staphylococcus identifying amplicons, a series of Staphylococcus genome segment sequences were obtained, aligned and scanned for regions where pairs of PCR primers would amplify products of about 45 to about 200 nucleotides in length and distinguish individual species, strains, and/or genotypes by their molecular masses or base compositions. A typical process shown in Figure 1 is employed for this type of analysis.

[00105] A database of expected base compositions for each primer region was generated using an in silico PCR search algorithm, such as (ePCR). An existing RNA structure search algorithm (Macke et al., Nucl. Acids Res., 2001, 29, 4724-4735, which is incorporated herein by reference in its entirety) has been modified to include PCR parameters such as hybridization conditions, mismatches, and thermodynamic calculations (SantaLucia, Proc. Natl. Acad. Sci. U.S.A., 1998, 95, 1460-1465, which is incorporated herein by reference in its entirety). This structure search algorithm can be used for other nucleic acids, such as DNA. This also provides information on primer specificity of the selected primer pairs.

[00106] Table 1 lists a collection of primers (sorted by primer pair number) configured to identify Staphylococcus bioagents using the methods described herein. The primer pair number is an inhouse database index number. Primer sites (conserved regions which primers were configured to hybridize within) were identified on Staphylococcus genes including arcC, aroE, ermA, ermC, gmk, gyrA, mecA, mecRl, mupR, nuc, pta, pvluk, tpi, tsst, tufB, and yqi. The forward and reverse primer names shown in Table 1 indicate the gene region of a bacterial genome to which the forward and reverse primers hybridize relative to a reference sequence. The forward primer name GYRA_NC002953-7005-9668_234_261_F indicates that the forward primer ("_F") hybridizes to the GyrA gene ("GYRA"), specifically to residues 234-261 ("234_261") of a reference sequence represented by a sequence extraction of coordinates 7005-9668 (SEQ ID NO.: 8) from GenBank gi number 49484912 (as indicated by cross-references in Table 2 for the prefix "GYRA_NC002953"). This sequence extraction reference includes sequence encoding for the gyrA gene ("GYRA"). The

primer pair name codes appearing in Table 1 are defined in Table 2. For example, Table 2 lists gene abbreviations and GenBank gi numbers that correspond with each primer name code. For example, for the above-mentioned primer pair has the code "GYRA_NC002953" and is thus configured to hybridize to sequence encoding the gyrA gene, and the extraction sequence (SEQ ID NO: 8) 7005-9668 corresponds to coordinates 7005-9668 of GenBank gi number 49484912, which is a Staphylococcus aureus sequence. One of skill in the art will understand how to determine the exact hybridization coordinates of the primers with respect to the GenBank sequences, given this information. The reference nomenclature in the primer name is selected to provide a reference, and does not necessarily mean that the primer pair has been configured with 100% complementarity to that target site on the reference sequence. One with ordinary skill knows how to obtain individual gene sequences or portions thereof from genomic sequences present in GenBank. In Table 1, Tp = 5-propynylcytosine; * = phosphorothioate linkage; 1 = inosine. T GenBank gi numbers for reference sequences of acteria are shown in Table 2 (below). In some cases, the reference sequences are extractions from bacterial genomic sequences or complements thereof. A description of the primer design is provided herein.

Table 1: Primer Pairs for Identification of Staphylococcus

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
258	RNASEP_SA_3 1_49_F	GAGGAAAGTCCAT GCTCAC	255	RNASEP_SA_338 _379_R	ATAAGCCATGTTC TGTTCCATC	312
258	RNASEP_SA_3 1_49_F	GAGGAAAGICCAI GCTCAC	255	RNASEP_EC_345 _362_R	ATAAGCCGGGTTC TGTCG	313
259	RNASEP_SA_3 1_49_b	GAGGAAAGTCCAT GCTCAC	255	RNASEP_BS_363 _384_R	GTAAGCCATGTTT TGTTCCATC	314
258	RNASEP_EC_6 1_77_F	GAGGAAAGICCGG GCTC	257	RNASEP_SA_358 _379_R	ATAAGCCATGTTC TGTTCCATC	312
258	RNASEP_EC_6 1_77_F	GAGGAAAGICCGG GCTC	257	RNASEP_EC_345 _362_R	ATAAGCCGGGTTC TGTCG	313

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
258	RNASEP_EC_6 1_77_F	GAGGAAAGICCGG GCTC	257	RNASEP_BS_363 384_R	GTAAGCCATGTTT TGTTCCATC	31.4
258	RNASEP_BS_4 3_61_F	GAGGAAAGTCCAT GCTCGC	256	RNASEP_SA_358 _379_R	ATAAGCCATGTTC TGTTCCATC	312
258	RNASEP_BS_4 3_61_F	GAGGAAAGICCAT GCTCGC	256	RNASEP_EC_345 _362_R	ATAAGCCGGGTTC TGTCG	313
258	RNASEP_BS_4 3_61_F	GAGGAAAGICCAT GCTCGC	256	RNASEP_BS_363 _384_R	GTAAGCCATGTTT TGTTCCATC	314
259	RNASEP_BS_4 3_61_F	GAGGAAAGTCCAT GCTCGC	256	RNASEP_BS_363 _384_R	GTAAGCCATGTTT TGTTCCATC	314
260	RNASEP_EC_6 1_77_F	CACCAAACTCCCC GCTC	257	RNASEP_EC_345 _362_R	ATAACCCOSCTTC TGTCG	313
262	RNASEP_SA_3 1_49_F	GAGGAAAGICCAT GCTCAC	255	RNASEP_SA_358 _379_R	ATAAGCCATGTTC TGTTCCATC	312
877	MEGA_Y14051 _3774_3802_ F	TAAAACAAACTAC GGTAACATTGATC GCA	57	MECA_Y14051_3 828_3854_R	TCCACATCTAACT TCCACATACCATC T	141
878	MECA_Y14051 _3645_3670_ F	TGAAGTAGAAATG ACIGAACGICCGA	56	MECA_Y14051_3 690_3719_R	TGATCCTGAATGT TTATATCTTTAAC GCCT	140
879	MEGA_Y14051 _4507_4530_	TCAGGTACTGCTA TCCACCCTCAA	58	MECA_Y14051_4 555_4581_R	TGGATAGACGTCA TATGAAGGTGTGC T	142
880	MECA_Y14051 _4510_4530_ F	TGTACTGCTATCC ACCCTCAA	59	MECA_Y14051_4 586_4610_R	TATTCTTCGTTAC TCATGCCATACA	143
881	MECA_Y14051 _4669_4698_ F	TCACCAGGTTCAA CTCAAAAAATATT AACA	61	MECA_Y14051_4 765_4793_R	TAACCACCCCAAG ATTTATCTTTTTG CCA	146
882	MEGA_Y14051 _4520_4530P _F	TCpCpACpCpCpT pCpAA	60	MECA_Y14051_4 590_4600P_R	TpACpTpCpATpG CpCpA	144
883	MEGA_Y14051 _4520_4530P _F	TCpCpACpCpCpT pCpAA	60	MECA_Y14051_4 600_4610P_R	TpATpTpCpTpTp CpGTpT	145
2056	MECI- R_NC003923- 41/96- 41609 33 60 _F	TTTACACATATOS TGAGCAATGAACT GA	62	MECI- R_NC003923- 41:98- 41609 86 113 R	TTGTGATATGGAG GTGTAGAAGGTGT TA	147
2057	AGR- III_NC00392 3-2108074- 2109507_1_2 3 F	TCACCAGTITGCC ACGIATCTICAA	191	AGR- III_NC003923- 2106074- 2109507_56_79 R	ACCTGCATCCCTA AACGTACTTGC	256
2058	AGR- III_NC00392 3-2108074- 2109507_569 _596_F	TGAGCTTTTAGTT CACITTTTCAACA GC	192	AGR- III_NC003923- 2108074- 2109507_622_6 53_R	TACTTCAGCTTCG TGCAATAAAAAAT CACAAT	267

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
2059	AGR- III_NC00392 3-2108074- 2109507_102 4_1052_F	TTTCACACAGCGT GTTTATAGTTCTA CCA	193	AGR- III_NC005923- 2108074- 2109507_1070_ 1098_R	TGTAGGCAAGTGC ATAAGAAATTGAT ACA	268
2060	AGR- I_AJ617706_ 622_651_F AGR-	TGGTGACTTCATA ATGGATGAAGTTG AAGT TGGGATTTTAAAA	217	AGR= AJ617706_69 4_726_R AGR=	TCCCCATTTAATA ATTCCACCTACTA TCACACT TGGTACTTCAACT	292
2061	I_AJ617706_ 580_611_F	AACATTGGTAACA TCGCAG	218	AJ617706_62 6_655_R	TCATCCATTATGA AGTC	293
2062	II_NCC02745 -2079448- 2080879_62C _651_F	TCTTGCAGCAGTT TATTTGATGAACC TAAAGT	219	II NG002745- 2079448- 2080879 700 7 31 R	TTGTTTATTGTTT CCATATGCTACAC ACTTTC	294
2063	11_NCC 02745 -2079448- 2080879_649 679_F	TGTACCCGCTGAA TTAACGAATTTAT ACGAC	220	_1_NC002745= 2079448- 2080879_715_7 45_R	TCGCCATAGCTAA GTTGTTTATTGTT TCCAT	1
2064	AGR- IV_AJ617711 931_961_F	TGGTGATAATGAC CTOGC	221	AGR- IV_AJ617711_1 004_1035_R	TECGCTATCAACG ATTTTGACAATAT ATGTGA	296
2065	AGR- IV_AJ617711 250_283_F	TGGCACTCITGCC TTTAATATTAGTA AACTATCA	222	AGR- IV_AJ617711_3 09_335_R	TCCCATACCTATG GCGATAACTGTCA T	297
2066	BLAZ_NC0029 52(1913827. .1914672)_6 8_68_F	TCCACTTATCGCA AATGGAAAATTAA GCAA	223	BLAZ_NC002952 (1913827191 4672)_68_68_R	TGGCCACTTTTAT CAGCAACCTTACA GTC	290
2067	BLAZ_NC0029 52(1913827. .1914672)_6 8_68_2_F	TGCACTTATCGCA AATGGAAAATTAA GCAA	224	BLAZ_NC002952 (1913827191 4672)_68_68_2 _R	TAGTCTTTTGGAA CACCGTCTTTAAT TAAAGT	291
2060	BLAZ_NC0029 52(1913827. .1914672)_6 0_60_3_F	TGATACTTCAACG CCTGCTGCITTC	225	BLAZ_NC002952 (191382791 4672)_68_68_3 _R	TGGAACACCGTCT TTAATTAAAGTAT CTCC	292
2069	BLAZ_NC0028 52(1913827. .1914672)_6 8_68_4_F	TATACTTCAACGC CTGCTGCTITC	226	BLAZ_NC002952 (1913827191 4672)_68_68_4 _R	TCTTTTCTTTGCT TAATTTTCCATTT GCGAT	293
2070	ELAZ_NC0029 52(1913827. .1914672)_1 _33_F	TGCAATTGCTTTA GTTTTAAGTGCAT GTAATTC	227	BLAZ_NC002952 (1913827191 4672) 34 67 R	TTACTTCCTTACC ACTTTTAGTATCT AAAGCATA	284
2071	52(1913827. .1914672)_3 _34_F	TCCITGCTITAGT TTTAAGTGCATGT AATICAA	228	BLAZ_NC002952 (191382791 4672) 40 68 R	TGGGGACTTCCTT ACCACTTTTAGTA TCTAA	285
2072	BSA- P_NCOC3923- 1304065- 1303589_99_ 125_F	TAGOGAATOTGGC TTTACTTCACAAT T	194	BSA- A_NC003923- 1304065- 1303509_165_1 93_R	TGCAAGGGAAACC TAGAATTACAAAC CCT	269

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
2073	BSA- A_NCOC3923- 1304065- 1303589_194 _218_F	ATCAATTTGGTGG CCAAGAACCTGG	195	BSA- A_NC003923- 1304065- 1303589_253_2 78_R	TGCATAGGGAAGG TAACACCATAGTT	270
2074	BSA- A_NC003923- 1304065- 1303589_328 349_F BSA- A_NC003923- 1304066- 1303589_253	TTGACTGCGGCAC ANCACGGAI TGCTATGGTGTTA	196	BSA- A NG003923- 1304065- 130589_388_4 15_R BSA- A NG003923- 1304065- 1303589_317_3	TAACAACGTTACC TTGGGGATCCACT AA TGTTGTGCCGCAG TCAAATATCTAAA	271
2075	278 F BSA- B_NC0C3923- 19:7149- 19:4156_953 982 F	CCTTCCCTATGCA TAGCAACAAATAT ATCTGAAGCAGCG TACT	197	44 R BSA- B_NC003923- 1917149- 1914156_1011_ 1039 R	TA TGTGAAGAACTTT CAAATCTGTGAAT CCA	272
2077	BSA- B_NC003923- 1917149- 1914156_105 C_1081_F	TCAAAACTATCCA TTTGAACAACTCG TGAATA	199	BSA- B_NC003923- 1917149- 1914156_1109_ 1136_R	TCTTCTTCAAAAA TTGTTGTCCCGAA AC	274
2078	BSA- B_NC0C3923- 1917149- 1914156_126 C_1286_F	TCATTATCATGCG CCAATGAGTGCAG A	200	BSA- B_NC003923- 1917149- 1914156_1323_ 1353_R	TGGACTAATAACA ATGAGCTCATTGT ACTGA	275
2079	B NC003923- 1917149- 1914156 212 6 2153 F ERMA NC0029	TTTCATCTIATCG AGGACCCGAAATC GA	201	B NC003923- 1317149- 1914156_2186_ 2216_R ERMA NC002952	TGAATATGTAATG CAAACCAGTCTTT GTCAT	276
2080	52-55890- 56621_366_3 92_F ERMA_NC0029 52-55890-	TOGCTATCTTATC GTTGAGAAGGGAT T TAGCTATCTTATC	28	-55890- 56621_487_513 R ERMA_NC002952 -55890-	TGAGTCTACACTT GGCTTAGGATGAA A TGAGCATTTTTAT	114
2081	56621_366_3 95 F ERMA NC0029	GTTGAGAAGGGAT TTCC	294	56621_438_465 R ERMA NCC02952	ATCCATCTCCACC AT	295
2082	52-55890- 56621_374_4 02_F	TGATCGTTGAGAA GGGATTTGCGAAA AGA	27	-55890- 56621_473_504 R	TCTTGGCTTAGGA TGAAAATATAGTG GTGGTA	113
2083	52-55890- 56621_404_4 27_F	TGCAAAATCTGCA ACGAGCTTTGG	29	=55890= =56621_491_520 _R	TCAATACAGAGTC TACACTTGGCTTA GGAT	115
2084	ERMA_NC0029 52=55890- 56621_489_5 16_F	TCATCCTAAGCCA AGTGTAGACTCTG TA	30	ERMA_NCC02952 -55890- 5662'_586_615 _R	TGGACGATATTCA CGGTTTACCCACT TATA	116

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
2085	ERMA_NC0029 52-55890- 56621_586_6 14_F	TATAAGTGGGTAA ACCGTGAATATOG TOT	31	-55890- 56621_640_665 R	TTGACATTTGCAT GCTTCAAAGCCTG	117
2086	ERMS_NC0059 08=2004= 2738_85_116 F	TOTGAACATGATA ATATOTTTGAAAT OGGOTO	35	ERMC_NCC05908 -2004- 2738_173_206_ R	TCCGTAGTTTTGC ATAATTTATGGTC TATTTCAA	121
2087	ERMC_NC0059 08-2004- 2738_90_120 F	TCATGATAATATC TTTGAAATCGGCT CAGGA	33	ERMC_NCC05908 -2004- 2738_16C_189_ R	TTTATGGTCTATT TCAATGGCAGTTA CGAA	119
2088	ERMC_NG0059 08-2004- 2738_1_5_13 9 F	TCAGGAAAAGGGC ATTTTACCCTTG	34	ERMC_NCC05908 -2004- 2738_16187_ R	TATGGTCTATTTC AATGGCAGTTACG A	120
2089	ERMC_NC0059 08=2004= 2738_374_39 7 F	TAATOGTGGAATA CGGGTTTGCTA	36	ERMC_NCC05908 -2004- 2738_425_452_ R	TCAACTTCTGCCA TTAAAAGTAATGC CA	122
2090	ERMC_NC0059 08-2004- 2738_101_12 5 F	TCTTTGAAATCGG CTCAGGAAAAGG	32	ERMC_NCC05908 -2004- 2738_159_108_ R	TGATGGTCTATTT CAATGGCAGTTAC GAAA	118
2091	ERMB_Y1360C -625- 1362_291_32 I_F	TGTTGGGAGTATT CCTTACCATTTAA GCACA	229	ERMB_Y13600- 625- 1362_352_380_ R	TCAACAATCAGAT AGATGTCAGACGC ATG	296
2092	ERMB_Y1360C -625- 1362_344_36 7_F	TGGAAAGCCATGC GTCTGACATCT	23C	ERMB_M13600- 625- _362_415_437_ R	TGCAAGAGCAACC CTAGTGTTCG	297
2093	ERMB_Y1360C -625- 1362_404_42 S_F	TGGATATTCACCG AACACTAGGGTTG	231	ERMB_Y13600= 625= 1362_471_493_ R	TAGGATGAAAGCA TTCCGCTGGC	298
2094	FRMB_Y1360C -625- 1362_465_40 7_F	TAAGCTGCCAGCG GAATGCTTIC	232	ERMB_Y13600- 625- 1362_521_545_ R	TCATCTGTGGTAT GGCGGGTAAGTT	259
2095	PVLUK_NC003 923- 1529595- 1531285_688 713 P	TGAGCTGCATCAA CTGTATTGGATAG	39	2VLUK NC00392 3-1529595- 1531285_775_8 C4_R	TGGAAAACTCATG AAATTAAAGTGAA AGGA	125
2096	PVLUK_NC003 923- 1529595- 1531285_103 9 1068 F	TSGAACAAAATAG TOTOTOSGATTTT GACT	37	PVLUK_NC00392 3-1529595- 1531285 1095 1125_R	TCATTAGGTAAAA TGTCTGGACATGA TCCAA	123
2097	PVLUK_NCO03 923- 1529595- 1531285_908 936_P	TGAGTAACATCCA TATTTCTGCCATA CGT	40	PVLUK NC00392 3-1529595- 1531285_950_9 78_R	TCTCATGAAAAAG GCTCAGGAGATAC AAG	126

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
2098	PVLUK_NC003 923- 1529595- 1531285_610 _633_F	TOGGAATCTGATG TTCCACTTCTT	38	">VIAIK_NC00392 3-1529595- 1531285 654 6 82_R	TCACACCTGTAAG TGAGAAAAAAGGTT GAT	124
2099	\$A442_NC003 923= 2538576= 2538831_11_ 35_F	TGTCGGTACACGA TATICTTCACGA	205	SA442_NC00392 3-2538576- 2538831_98_12 4_R	TTTCCGATGCAAC GTAATGAGATTTC A	13
2100	SA442_NC003 923= 2538576- 2538831_98_ 124_F	TGAAATCTCATTA CGTIGCATCGGAA A	206	SA442 NC00392 3-2538576- 2538831_163_1 88_R	TCGTATGACCAGC TTCGGTACTACTA	14
2101	SA442_NC003 923- 2538576- 2538831_103 126 F	TCTCATTACGTTG CATCGGAAACA	207	SA442_NC00392 3-2538576- 2538831_161_1 87 R	TTTATGACCAGCT TCGGTACTACTAA A	15
2102	SA442_NC003 923- 2538576- 2538831_166 _188_F SEA_NC00392	TAGTACCGAAGCT GGTCATACGA	208	SA442_NC00392 3-2538576- 2536831_231_2 57_R SEA_NC003923-	TCATAATCAACCC AAACCTTTTTCAC G	96
2103	3=2052219= 2051456_115 135_F	TGCAGGGAACAGC TTTAGGCA	209	2052219- 2051456_173_2 00_R	TCGATCGTGACTC TCTTTATTTTCAG TT	91
2104	SEA_NC00392 3-2052219- 2051456_5/2 _598_F	TAACTCTGATGTT TTTGATGGGAAGG T	210	SEA_NC003923- 2052219- 2051456_621_6 51_R	TGTAATTAACCGA AGGTTCTGTAGAA GTATG	98
2105	SEA_NC00392 3-2052219- 2051456_382 _414_F	TGTATGGTGGTGT AACGTTACATGAT AATAATC	211	SEA_NC003923- 2052219- 2051456_464_4 92_B	TAACCGTTTCCAA AGGTACTGTATTT TGT	317
2106	SEA_NC00392 3-2052219- 2051456_377 _406_F	TTGTATGTATGGT GGTGTAACGTTAC ATGA	212	SEA_NC003923- 2052219- 2051456_459_4 92_R	TAACCGTTTCCAA AGGTACTGTATTT TGTTTACC	318
210/	SEB_NC00275 8-2135540- 2135140_208 _23/_F	TTTCACATGTAAT TTTGATATICGCA CTGA	247	SEB_NC002758- 2135540- 2135140_273_2 98_R	TCATCTGGTTTAG GATCTGGTTGACT	394
2108	SEB_NC00275 8=2135540= 2135140_206 235 F	TATITCACATGTA ATTITGATATTCG CACI	248	SEB_NC002750- 2135540- 2135140_281_3 04_R	TGCAACTCATCTG GTTTAGGATCT	305
2109	SEB_NC00275 8-2135540- 2135140_402 _402_F	TARCARCTCGCCT TATGAAACGGGAT ATA	249	SEB_NC002758- 2135540- 2135140_402_4 02_R	TGTGCAGGCATCA TGTCATACCAA	306

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
2110	SEB_NC00275 8=2135540= 2135140_402 402_2_F SEC_NC00392	TTGTATGTATGGT GGTGTAACTGAGC A	25C	SEB NC002758- 2135540- 2135140 402 4 02 2 R SEC NC005923-	TTACCATCTTCAA ATACCCGAACAGT AA	307
2111	3-851678- 852768_546_ 575 F	TTAACATGAAGGA AACCACTTTGATA ATGG	213	851678- 852768_620_64 7_R	TGAGTTTGCACTT CAAAAGAAATTGT GT	319
2112	SEC_NC00392 3-851678- 852768_537_ 566_F SEC_NC00392	TGGAATAACAAAA CATGAAGGAAACC ACTI	214	SEC_NC003923= 051670= 852768_619_64 7_R SEC_NC003923=	TCAGTTTGCACTT CAAAAGAAATTGT GTT	320
2113	3-851678- 852768_720_ 749 F	TGACTTTAACAGT TCACCATAIGAAA CAGG	215	851678- 852768_794_81 5 R	TCGCCTGGTGCAG GCATCATAT	321
2114	SEC_NC00392 3-851678- 852768_787_ 010_F	TGGTATGATATGA TGCCTGCACCA	216	SEC_NC003923- 851678- 852768_853_88 6_R	TCTTCACACTTTT AGAATCAACCGTT TTATTGTC	322
2115	SED_M28521_ 657_602_F	TGGIGGTGAAATA GATAGGACIGCTT	183	SED_M28521_74 1_770_R	TGTACACCATTTA TCCACAAATTGAT TGGT	258
2116	SED_M28521_ 690_711_F	TGGAGGTGTCACT CCACACGAA	184	SED_M28521_73 9_770_R	TGGGCACCATTTA TCCACAAATTGAT TGGTAT	259
2117	SED_M28521_ 833_854_F	TTGCACAAGCAAG GCGCTATTI	185	SED_M28521_88 8_911_R	TCGCGCTGTATTT TTCCTCCGAGA TGTCAATATGAAG	260
2118	SED_M28521_ 962_98/_F SEA-	TGGATGTTAAGGG TGATTTTCCCGAA	186	SED_M28521_10 22_1048_R SEA-	GTGCTCTGTGGAT A	251
2119	SEE_NC00295 2-2131289- 2130703_16_ 45_F	TTTACACTACTTT TATTCATTGCCCT AACS	233	SEE_NC002952- 2131289- 2130703_71_98 _R	TCATTTATTTCTT CGCTTTTCTCGCT AC	290
2120	SEA- SEE_NC00295 2-2131209- 2130703_249 _2/8_F	TGATCATCCGTGG TATAACGAITTAT TAGI	234	SEA- SEE NC002952- 2131209- 2130703_314_3 44_R	TAAGCACCATATA AGTCTACTTTTT CCCTT	291
2121	SEE_NC00295 2-2131289- 2130/03_409 _437_F	TGACATGATAATA ACDEATTGACCGA AGA	235	SEE_NC002952- 2131289- 2130703_465_4 94_R	TCTATAGGTACTG TAGTTTGTTTTCC GTCT	323
2122	SBE_NC00295 2=2131289= 2130703_525 550 F	TGTTCAAGAGCTA GATCTTCACGCAA	236	SEE_NC002952- 2131289- 2130703_586_5 86_R	TTTGCACCTTACC GCCAAAGCT	324
2123	SEE_NC00295 2-2131289- 2130703_525 _549_F	TGTTCAAGAGCTA GATCTTCAGGCA	237	SEE NC002952~ 2131289- 2130703_506_5 86_2_R	TACCTTACCGCCA AAGCTGTCT	325

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
2124	SEE_NC00295 2-2131289- 2130703_361 384_F SEG_NC00275	TCTGGAGGCACAC CAAATAAAACA	238	SEE NC002952- 2131289- 2130703 444 4 71 R SEG NC002758-	TCCGTCTATCCAC AAGTTAATTGGTA CT	326
2125	B=1955100= 1954171_225 _251_F SEG_NC00275	TGCTCAACCCGAT CCTAAATTAGACG A	251	1955100= 1954171_321_3 46_R SEG NC002758=	TAACTCCTCTTCC TTCAACAGGTGGA	308
2126	0-1955100- 1954171_623 651_F SEG_NC0027E	TGGACAATAGACA ATCACTTGGATTT ACA	252	1955100- 1954171_671_7 02_k SEG_NC002758-	TGCTTTGTAATCT AGTTCCTGAATAG TAACCA	309
2127	B-1955100- 1954171_540 _564_F SEG_NC00275	TGGAGGTTGTTGT ATGTATGGTGGT	253	1955100- _954171_607_6 35_R SEG NC002758-	TGTCTATTGTCGA TTGTTACCTGTAC AGT	310
2126	8955100- 1954171_694 _710_F SEH_NC00295	TACANAGCAAGAC ACTGGCTCACTA	254	_955100= _954171_735_7 62_F SEH NC002953=	TGATTCAAATGCA GAACCATCAAACT CG	311
2129	3-60024- 60977_449_4 72_F SEH NC00295	TTGCAACTGCTGA TTTAGCTCAGA	239	6002(- 60977_547_576 _R SEH NC002953-	TAGTGTTGTACCT CCATATAGACATT CAGA	327
2130	3-60024- 60977_408_4 34_s' SEII NC00295	TAGAAATCAAGGT GATAGTGGCAATG A	24C	60024- 60977_450_473 _R SEII NC002953-	TTCTGAGCTAAAT CAGCAGTTGCA	328
2131	3-60024- 60977_547_5 76_F SEH_NC00295	TCTGAATGICTAT ATGGAGGTACAAC ACTA	241	60024- 60977_608_634 R SEH NC002953-	TACCATCTACCCA AACATTAGCACCA A	298
2132	3-60024- 60977_546_5 75_3 SRT_VC00275	TYCIGAATGICYA TATGGAGGIACAA CACI	242	60024- 60977_594_616 _R SET_NC002758-	TAGCACCAATCAC CCTTTCCTGT	299
2133	B=1957830= 1956949_324 _349_F SET_NC00275	TCAACTCGAATTT TCAACAGGTACCA	243	1957830- 1956949_419_4 46_R SEI NC002758-	TCACAAGGACCAT TATAATCAATGCC AA	300
2134	8-1957830- 1956949_336 _363_F	TTCAACAGGTACC AATGATTTGATCT CA	244	1957830- 1956949_420_4 47_k	TGTACAAGGACCA TTATAATCAATGC CA	301
2135	SEI_NC00275 B-1957830- 1956949_356 _384_F	TGATCTCAGAATC TAATAATTGGGAC GAA	245	SEI_NC002758- 1957830- _956949_449_4 74_R	TCTGGCCCCTCCA TACATGTATTTAG	302
2136	SEI_NC00275 8=_957830= 1956949_223 _253_F	TCTCAAGGIGATA TTGGTGTAGGTAA CTTAA	246	SEI_NC002758- _957830- _956949_290_3 _16_R	TGGGTAGGTTTTT ATCTGTGACGCCT T	303
2137	SEJ_AF05314 C_1307_1332 F	TGTGGAGTAACAC TGCATGAAAACAA	187	SEJ_AF053140_ 1301_1404_R	TCTAGCGGAACAA CAGTTCTGATG	262

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
2136	SEJ_AF05314 C_13/8_1403 F	TAGCATCAGAACT GTTGTTCCGCTAG	188	SEJ_AF053140_ 1429_1458_R	TCCTGAAGATCTA GTTCTTGAATGGT TACT	263
2139	SEJ_AF05314 (431459 F SEJ_AF05314	TAACCATTCAGA ACTAGATCITCAG GCA TCATTCAAGAACT	189	SEJ_AF053140_ 1300_1331_R	TAGTCCTTTCTGA ATTTTACCATCAA AGGTAC TCAGGTATGAAAC	264
2140	C_1434_1461 F TSST NC0027	AGATCTTCAGGCA AG	190	SEJ_AF053140_ 1521_1549_R TSST_NC002758	ACGATTAGTCCTT TCT	265
2141	58-2137564- 2138293_206 236_F	TOCTTAGATAAT TOCTTAGGATCTA TOCCT	66	-2137564- 2138293_278_3 C5_R	TGTAAAAGCAGGG CTATAATAAGGAC TC	151
2142	TSST_NC0027 58=2137564= 2138293_232 _258_F	TGCGTATAAAAA CACAGATGGCAGC A	67	TSST_NCC02758 -2137564- 2138293_289_3 13_R	TGCCCTTTTGTAA AACCACCCCTAT	152
2143	TSST_NC0027 58-2137564- 2138293_382 410 F	TOCAAATAAGTGG OGTTACAAATACT GAA	68	TSST_NCC02758 -2137564- 2138293_448_4 78_R	TACTTTAAGGGGC TATCTTTACCATG AACCT	153
2144	TSST_MC0027 58-2137564- 2138293_297 _325_F	TCTTTTACAAAAG GGGAAAAAGTTGA CTT	69	TSST_NCC02758 -2137564- 2138293_347_3 73_R	TAAGTTCCTTCGC TAGTATGTTGGCT T	154
2145	ARCC_NC0039 23-2725050- 2724595_37_ 58 F	TOGOOGGCAATGC CATTGGATA	75	ARCC_NCC03923 -2725050- 272(595_97_12 8 R	TGAGTTAAAATGC GATTGATTTCAGT TTCCAA	161
2146	ARCC_NC0039 23-2725050- 2724595_131 161 F	TGAATAGTGATAG AACTGTAGGCACA ATOGT	72	ARCC_NC003923 -2725050- 2724595_214_2 45 R	TCTTCTTCTTTCG TATAAAAAGGACC AATTGG	156
2147	ARCC_NC0039 23-2725050- 2724595_218 249 F	TTGGTCCTTTTA TACGAAAGAAGAA GTTGAA	74	ARCC_NCC03923 -2725050- 2724595_322_3 53 R	TGGTGTTCTAGTA TAGATTGAGGTAG TGGTGA	160
2148	AROE NC0039 23-1674726- 1674277_371 393 F	TTGCGAATAGAAC GATGGCTCGT	80	AROE NCC03923 -1674726- 1674277_435_4 66 R	TCGAATTCAGCTA AATACTTTTCAGC ATCT	167
2149	AROE_NC0039 23-1674726- 1674277_30_ 62_F	TGGGGCTTTAAAT ATTCCAATTGAAG ATTTTCA	79	AROE_NCC03923 -1674726- 1674277_155_1 B1_R	TACCTGCATTAAT CGCTTGTTCATCA A	166
2150	AROP_NC0039 23-1674726- 1674277_204 _232_F	TGATGGCAAGTGG ATAGGGTATAATA CAG	7.6	AROR_NCC03923 -1674726- 1674277_308_3 35_R	TAAGCAATACCTT TACTTGCACCACC TG	162
7151	GLPF_NC0039 23-1296927- 1297391_27C 301_F	TGCACCGGCTATT AAGAATTACTTTG CCAPCT	202	GLPF_NC003923 -1296927- 1297391_382_4 -4_R	TGCAACAATTAAT GCTCCGACAATTA AAGGATT	277

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
2152	GLPF_NC0039 23-129692/- 1297391_27_ 51_F	TGGATGGGGATTA GCGGTTACAATG	203	GLPF_NC003923 -129692/- 1297391_81_10 8_R	TAAAGACACCGCT GGGTTTAAATGTG CA	278
2153	GLP: NC0039 23-1296927- 1297391_239 _260_F	TAGCTGGGGGGAA ATTAGGTGI	204	GLPE_NC003923 -1296927- 1297391_323_3 59_R	TCACGGATAAATA AAATACCTAAAGT TAATGCCATTG	279
2154	GMK_NC00392 3-1190906- 1191334_91_ 122_F	TACTTTTTTAAAA CTAGGGATGCGTT TGAAGC	81	GMK_NC003923- 1190906- 1191334_166_1 97_R	TGATATTGAACTG GTGTACCATAATA GTTGCC	168
2155	GMK_NG00392 3-1190906- 1_91334_240 _267_F	TGAAGTAGAAGGT GCAAAGCAAGTTA GA	82	GMK_NC003923- 1190906- 1191334_305_3 33_R	TCGCTCTCTCAAG TGATCTAAACTTG GAG	169
2156	GMK_NC00392 3-1_90906- 1191334_301 _329_F	TCACCTCCAAGTT TAGATCACTTGAG AGA	83	GMK_NC0C3923- 1190906- 1191334_403_4 32_R	TGGGACGTAATCG TATAAATTGATCA TTTC	170
2157	PUA_NC00392 3-628885- 629355_237_ 263 F	TCTTGTTTATGCT GGTAAAGCAGATG G	87	PIA_NC003923- 628885- 629355_314_34 5 R	TGGTACACCTGGT TTCGTTTTGATGA TTTGTA	172
2156	PTA_NC00392 3-620805- 629355_141_ 171 F	TGAATTAGTTGAA TCATTTGTTGAAC GACGT	84	PTA_NC0C3923- 620005- 629355_211_23 9 R	TGCATTGTACCGA AGTAGTTCACATT GTT	171
2159	PTA_NC00392 3-628885- 629355_328_ 356 F	TCCAAACCAGGTG TATCAAGAACATC AGG	88	PTA_NC0C3923- 628885- 629355_393_42 2 R	TGTTCTGGATTGA TTGGACAATCACC AAAG	175
2160	TPI_NC00392 3-830671- 831072_131_ 160 F	TGCAAGTTAAGAA AGCTGTTGCAGGT TTAT	89	TPI_NC003923= B30671= 831072_209_23 9 R	TGAGATGTTGATG ATTTACCAGTTCC CATTC	1/6
2161	TPT_NG00392 3=830671= 831072_1_34 F	TCCCACGAAACAG ATGAAGAAATTAA CAAAAAAG	90	TPT_NC0C3923- 830671- 031072_97_129 R	TGGTACAACATCG TTAGCTTTACCAC TTTCAGG	177
2162	TPI_NC00392 3-830671- 831072_199_ 227 F	TCAAACTGGGCAA TCGGAACTGGTAA ATC	91	TPI_NC0C3923= 830671- 831072_253_28 6 R	TGCGAGCAATAGT TTGACGTAGAAAT GCAGACAT	178
2163	YQI_NC00392 3-378916- 379431_142_ 167 F	TGAATTGCTGCTA TGAAAGGTGGCTT	93	YQI_NC003923- 378916- 379431_259_28 4 R	TCGCCAGCTAGCA CGATGTCATTTTC	190
2164	YQI_NC00392 3-378916- 379431_44_7 7_F	TACAACATATTAT TAAAGAGACGGGT TTGAATCC	95	YQI_NC0C3923- 378916- 379/31_120_14 5_R	TTCGTGCTGGATT TTGTCCTTGTCCT	182
2165	YQT_NC00392 3-378916- 379431_135_ 160_F	TOCAGCACGAATT GCTGCTATGAAAG	92	YQT_NC0C3923= 378916= 379431_193_22 1_R	TCCAACCCAGAAC CACATACTTTATT CAC	179

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
2166	YQI_NC00392 3-3/8916- 379431_275_ 300_F	TAXTGXTGGTAT GGAGAATATGTCT	94	YQI_NC003923- 3/8916- 379431_364_39 6_R	TCCATCTGTTAAA CCATCATATACCA TCCTATC	181
2167	BLAZ_(19138 271914672) 546_575_F BLAZ_(19138	TOCACTTATCGCA AATGGAAAATTAA GCAA	223	BLAZ_(1913827 1914672)_65 5_663_R	TGGCCACTTTAT CAGCAACCTTACA GTC	230
2168	271914672)_546_575_2 F BLAZ_(19138	TGCACTTATCGCA AATGGAAAATTAA GCAA	224	BLAZ_(1913827 1914672)_62 8_659_R BLAZ_(1913827	TAGTCTTTTGGAA CACCGTCTTTAAT TAAAGT TGGAACACCGTCT	291
2169	271914672) 507 531 F BLAZ (19138	TGATACTTCAACG CCTGCTGCTTTC	225	1914672)_62 2_651_R BLAZ (1913827	TTAATTAAAGTAT CTCC TCTTTTCTTTGCT	292
2170	271914672) 508 531 F BLAZ_(19138	TATACTTCAACGC CTGCTGCTITC TGCAATTGCTTTA	226	1914672)_55 3_583_R BLAZ_(1913827	TAATTTTCCATTT GCGAT TTACTTCCTTACC	293
2171	27IS14672)_24_56_F BLAZ_(19138	GTTTTAAGTGCAT GTAATTC TCCTTGCTTTAGT	227	1914672)_12 154_R BLAZ_(1913827	ACTTTTAGTATCT AAAGCATA TGGGGACTTCCTT	294
2172	271S14672)_26_58_F BLAZ_NC0029 52-1913827-	TTTAAGTGCATGT AATICAA TCCACTTAICCCA	228	1914672)_12 7_157_R BLAZ_NC002952 -1913827-	ACCACTTTTAGTA TCTAA TCCCCACTTTTAT	295
2173	1914672_546 575 F BLAZ NC0029	AATGGAAAATTAA GCAA	223	1914672_655_6 83_K BLAZ NC002952	CAGCAACCTTACA GTC	290
2174	52-1913827- 19:4672_546 _575_2_F	TGCACTTATCGCA AATGGAAAATTAA GCAA	224	-1913827- :914672_628_6 59_R	TAGTCTTTTGGAA CAGCGTCTTTAAT TAAAGT	291
2175	BLAZ_NC0029 52-1913827- 1914672_507 _531_F	TGATACTTCAACG CCTGCTGCTTTC	225	BLAZ_NC002932 -1913827- 1914672_622_6 51_R	TGGAACACCGTCT TTAATTAAAGTAT CTCC	292
2176	BLAZ_NC0029 52-1913827- 1914672 508 531_F	TATACTTCAACGC CTGCTGCTITC	226	BLAZ_NC002952 -1913827- 1914672 553 5 83_R	TCTTTTCTTTGCT TAATTTTCCATTT GCGAT	293
2177	BLAZ_NC0029 52-1913827- 1914672_24_ 56_F	TCCAATTOCTTTA GTTTTAAGIGCAT GTAATTC	927	BLAZ_NC002952 -1913827- 1914672_121_1 54_R	TTACTTCCTTACC ACTTTTAGTATCT RAAGCATA	284
2178	BLAZ NC0029 52-1913827- 19'4672_26_ 58 F	TCCTTGCTTTAGT TTTAAGTGCATGT AATICAA	228	BLAZ NC002952 -1913827- '914672_127_1 57 R	TGGGGACTTCCTT ACCACTTTTAGTA TCTAA	295
2247	TUFB_NC0027 58-615038- 616222_693_ 721_F	TGTTGAACGTGGT CAAATCAAAGTTG GTG	46	TUFB_NCC02758 -615038- 616222_793_82 0 R	TGTCACCAGCTTC AGCGTAGTCTAAT ///	132
2248	TUFB_NC0027 58-615038- 616222 690 716 F	TOSTGTTGAACGT GGTCAAATCAAAG T	45	TUFB_NCC02758 -615038- 616222 793 82 0 R	TGTCACCAGCTTC AGCGTAGTCTAAT AA	132

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
2249	TUFB_NC0027 58-615038- 616222_696_ 725_F	TGAACGTGGTCAA ATCAAAGTTGGTG AAGA	47	TUFB_NCC02758 -615038- 616222_793_82 0_R	TGTCACCAGCTTC AGCGTAGTGTAAT AA	132
2250	TUFB_NC0027 58-615038- 616222_486_ 513_F	TOCCAGGTGACGA TGTACCTGTAATC	42	-615038- -616222_601_63 0_R	TGGTTTGTGAGAA TCAGGTTCTGGAG TTGG	128
2251	TUFB_NC0027 56-615038- 616222_945_ 972_F	TGNAGGTGGACGT CACACTCCATTCT TC	51	TUFB_NCC02758 -615038- 616222_1030_1 060_R	TAGGCATAACCAT TTCAGTACGTTCT GGTAA	135
2252	TUFB_NC0027 58-615038- 6_6222_333_ 356 F	TOCAATGCCACAA ACTCGTGAACA	41	TUFD_NCC02758 -615038- 616222_424_45 9 R	TTCGATTTGAACT AATTCTAATAATT CTTCATCGTG	127
2253	NUC_NC00275 8=894288= 894974_402_ 424_F	TOCTGAAGCAAGT GCATTTACGA	52	NUC_NC0C2758- 894288- 894974_483_5C 9 R	TACGCTAAGCCAC GTCGATATTTATC A	136
2254	NUC_AC00275 8-894288- 894974_53_8 1 F	TOCTTATAGGGAT GGCTATCAGTAAT GTT	53	NUC_NC0C2758- 894288- 894974_165_10 9 R	TGTTTGTGATGCA TTTGCTGAGCTA	137
2255	NUC_NC00275 8-094280- 894974_169_ 194_F	TCAGCAAATGCAT CACAAACAGATAA	54	NUC_NC0C2758- 894288- 894974_222_25 0_R	TAGTTGAAGTTGC ACTATATAGTGTT GGA	138
2256	NUC_NC00275 8-894288- 894974_316_ 345_F	TACAAAGGTCAAC CAATGACATTCAG ACTA	55	NUC_NC0C2758- 894288- 894974_396_42 1_R	TAAATGCAGTTGC TTCAGGGCCATAT	139
2309	MUPR_X75439 658689_ F	TOOTTTGATATAT TAIGCGATGGAAG GTTGGT	18	MUPR_X75439_1 /44_1//3_R	TCCGTTCCTTAAT ATGAGAAGGAAAC CACT	1.01
2310	MUPR_X75439 _'330_'353_ P	TTCCTCCTTTTGA AAGCGACGGTT	17	MUPR_X75439_1 413_1441_R	TGAGCTGGTGCTA TATGAACAATACC AGT	1.00
2312	MUPR_X75439 _1314_1338_ F	TTTCCTCCTTTTG AAAGCGACGGTT	16	MUPR_X75439_1 381_1409_R	TATATGAAGAATA CCAGTTCCTTCTG AGT	99
2313	MUPR_X73439 _2486_2516_ P	TAATTGGGCTCTT TCTCGCTTAAACA CCTTA	21	MUPR_X75439_2 548_2574_R	TTAATCTGGCTGC GGAAGTGAAATCG T	104
2314	MUPR_X73439 _2547_2572_ P	TACGATTTCACTT ODGCAGDCAGATT	23	MUPR_X75439_2 605_2630_R	TCGTCCTCTCGAA TCTCCGATATACC	109
2315	MUPR_X73439 _2666_2696_ P	TGOGTACAATACG CTTTATGAAATTT TAACA	24	MUPR_X75439_2 711_2740_R	TCAGATATAAATG GAACAAATGGAGC CACT	110
2316	MUPR_X75439 _2813_2843_ P	TAATCAAGCATTG GAAGATGAAATGC ATACC	25	MUPR_X75439_2 867_2890_R	TCTGCATTTTTGC GAGCCTGTCTA	111
2317	MUPR_X75439 _884_914_F	TGACATGGACTOC COCTATATAACTC TTGAG	26	MUPR_X75439_9 77_1007_R	TGTACAATAAGGA GTCACCTTATGTC CCTTA	112

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
2504	ARCC_NC0039 23-2725050- 2724595_135 _161P_F	TAGTPGATPAGAA CPTPGTAGGCPAC PAATPCPGT	73	ARCC_NC003923 -2725050- 2724595_214_2 39P_R	TCpTpTpTpCpGT ATAAAAAGGACpC pAATpTpGG	159
2505	PUA_NC00392 3=628885= 629355_237_ 263P_F	TCTTGTpTpTpAT GCbTpGGTAAAGC AGATGG	86	21A_NC003923- 628885- 629355_314_34 2P_R	TACpACpCpTGGT pTpTpCpGTpTpT pTpGATGATpTpT pGTA	174
2738	GYRA_NC0029 53-7005- 9668_166_19 5_F	TANGGTATGACAC CEGATAAATCATA TAAA	2	GYRA_NCC02953 -7005- 9668_265_287_ R	TCTTGAGCCATAC GTACCATTGC	5
2739	GYRA_NC0029 53-7005- 9668_221_24 9_F	TAATGGGTAAATA TCACCCTCATGGT GAC	3	GYRA_NCC02953 -7005- 9668_316_343_ R	TATCCATTGAACC AAAGTTACCTTGG CC	6
2740	GYRA_NC0029 53=7005= 9668_221_24 9_F	TAATGGGTAAATA TCACCCTCATGGT CAC	3	GYRA_NCC02953 -7005- 9668_253_283_ R	TAGCCATACGTAC CATTGCTTCATAA ATACA	7
2741	GYRA_NC0029 53-7005- 9660_234_26 1_F	TCACCCTCATGGT GACTCATCTATTT AT	4	GYRA_NCC02953 -7005- 9660_265_207_ R	TCTTGAGCCATAC GTACCATTGC	5
3004	TUFB_NC0027 58-615038- 616222_684_ 704_F	TACAGGCCGTGTT GAACGTGG	43	TUFB_NCC02758 -615030- 616222_778_80 9_R	TCACCCTACTCTA ATAATTTACGGAA CATTTC	129
3005	TUFB_NC0027 56-615038- 6_6222_688_ 710_F	TGCCGTGTTGAAC GTGGTCAAAT	44	TUFB_NCC02758 -615038- 616222_783_81 3_R	TGCTTCAGCGTAG TCTAATAATTTAC GGAAC	130
3006	TUFB_NC0027 58-615038- 616222_700_ 726_F	TGTGGTCAAATCA AAGTTGGTGAAGA A	49	TUFB_NCC02758 -615038- 616222_778_80 7_R	TGCGTAGTCTAAT AATTTACGGAACA TTTC	134
3007	TUFB_NC0027 58-615038- 616222_702_ 726_F	TGGTCAAATCAAA GTTGGTGAAGAA	50	TUFB_NCC02758 -615038- 616222_778_00 7_R	TGCGTAGTCTAAT AATTTACGGAACA TTTC	134
3008	TUFB_NC0027 58-615038- 616222_696_ 726_F	TCAACCTCCTCAA ATCAAAGTTGGTG AAGAA	48	TUFB_NCC02758 -615038- 616222_785_81 8_R	TCACCACCTTCAC CGTAGTCTAATAA TTTACGGA	133
3009	TUFB_NC0027 56-615038- 6_6222_690_ 716_F	TOGTGTTGAACGT GGTCAAATCAAAG T	45	TUFB_NCC02758 -615038- 616222_778_81 2_R	TCTTCAGCGTAGT CTAATAATTTACG GAACATTTC	131
3010	MECT- R_NC003923- 7.1796- 41609_36_59 _F	TCACATATCGTGA GC/ATG/ACTG	63	MECI- R_NC003923= 41798- 41609_09_112_ R	TGTGATATGGAGG TGTAGAAGGTG	148

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
3011	MECI- R_NC003925- 41798- 41609_40_66 _F	TGGGCGTGAGCAA TGAACTGATTATA C	64	MECI- R_NC003923- 41798- 41609_01_10_ R	TGGGATGGAGGTG TAGAAGGTGTTAT CATC	149
3012	MECI- R_NC003923- 41798- 41609_33_60 _2_F	TGGACACATATOG TGAGCAATGAACT GA	62	MECI- R_NC003923- 41798- 41609_81_110_ R	TGGGATGGAGGTG TAGAAGGTGTTAT CATC	149
3013	MECI- R_NC003923- 41798- 41609_29_60 _F	TGGGTTTACACAT ATCGTGAGCAATG AACTGA	65	MECI- R_NC003923- 41798- 41609_81_113_ R	TGGGGATATGGAG GTGTAGAAGGTGT TATCATC	150
3014	MUPR_X75439 _2490_2514_ F	TOCCCTCTTTCTC GCTTAAACACCT	20	MUPR X75439 2 548_2570_R	TCTCCCTCCCCAA GTGAAATCGT	103
3015	MUPR_X75439 _2490_2513_ F	TGGGCTCTTTCTC GCTTAAACACC	19	MUPR_X75439_2 547_2568_R	TGGCTGCGGAAGT GAAATCGTA	102
3016	MUPR_X75439 _2482_2510_ F	TAGATAATTGGGC TCTTTCTCGCTTA AAC	22	MUPR_X75439_2 551_2573_R	TAATCTGGCTGCG GAAGTGAAAT	106
3017	MUPR_X75439 _2490_2514_ F	TGGGCTCTTTCTC GCTTAAACACCT	20	MUPR_X75439_2 549_2573_R	TAATCTGGCTGCG GAAGTGAAATCG	105
3018	MUPR_X75439 _2482_2510_ F	TAGATAATTGGGC TCTTTCTCGCTTA AAC	22	MUPR_X75439_2 559_2589_R	TGGTATATTCGTT AATTAATCTGGCT GCGGA	138
3019	MUPR_X75439 _2490_2514_ F	TGGGCTCTTTCTC GCTTAAACACCT	20	MUPR_X75439_2 554_2501_R	TCGTTAATTAATC TCCCTCCCCAACT GA	107
3020	AROE_NC0039 23-1674726- 1674277_204 _232_F	TGATGGCAAGTGG ATAGGGTATAATA CAG	76	AROE_NC003923 -1674726- 1674277_309_3 35_R	TAAGCAATACCTT TACTTGCACCACC T	163
3021	AROE NC0039 23-1674726- 1674277_207 232 F	TGGCGAGTGGATA GGGTATAATACAG	78	AROE NCC03923 -1674726- 1674277_311_3 39_R	TTCATAAGCAATA CCTTTACTTGCAC CAC	165
3022	AROE_NC0039 23-1674726- 1674277_207 232P_F	TGGCpAAGTpGGA TpAGGGTpATpAA TpACpAG	77	AROE_NC003923 -1674726- 1674277_311_3 35P_R	TAAGCAATACCPT pTpTpACTpTpGC pACpCpAC	164
3023	ARCC_NC0039 23-2725050- 2724595_124 155_F ARCC_NC0039	TCTGAAATGAATA GTGATAGAACTGT AGGCAC	71	ARCC_NCC03923 -2725050- 2724595_214_2 45_R	TCTTCTTCTTTCG TATAAAAAGGACC AATTGG	156
3024	ARCC_NC0039 23-2725050- 2724595_131 61_F	TGANTAGTGATAG AACTGTAGGCACA ATOGT	72	ARCC_NC003923 -2725050- 2724595_212_2 42_R	TCTTCTTTCGTAT AAAAAGGACCAAT TGGTT	157

Primer Pair Number	Forward Primer Name	Forward Sequence	Forward SEQ ID NO.	Reverse Primer Name	Reverse Sequence	Reverse SEQ ID NO.
3025	ARCS_NC0039 23-2725050- 2724595_131 61_F	TGAATAGTGATAG AACTGTAGGCACA ATOCT	72	ARCC_NC003923 -2/25050- 2724595_232_2 60_R	TGCGCTAATTCTT CAACTTCTTCTTT CCT	158
3026	P.VA_NC00392 3-628885- 629355_231_ 259_F	TACAATGCTTGTT TATCCTCCTAAAC CAG	85	21A_NC0C3923- 628885- 629355_322_35 1_R	TGTTCTTGATACA CCTCCTTTCCTTT TGAT	173
3027	PTA_NC00392 3-628885- 629355_231_ 259_F	TACAATGCTTGTT TATGCTGGTAAAG CAG	85	PTA_NC0C3923= 628885= 629355_314_34 5_R	TGGTACACCTGGT TTCGTTTTGATGA TTTGTA	172
3028	PTA_NC00392 3-628885- 629355_237_ 263 F	TCTTGTTTATGCT GGTAAAGCAGATG G	87	PTA_NC0C3923- 628885- 629355_322_35 1 R	TGTTCTTGATACA CCTGGTTTCGTTT TGAT	173
3105	TSST1_NC002 758.2_35_57 _F	TAAGCCCTTTGTT GCTTGCGACA	329	TSST1_NC00275 8.2_146_1/3_R	TCAGACCCACTAC TATACCAGTCTAG CA	330
3106	758.2- 2137509- 2130213 519 546 F	TOSTCATCAGCTA ACTCAAATACATG GA	70	TSST1_NC00275 8.2-2137509- 2138213 593- 620_R	TCACTTTGATATG TGGATCCGTCATT CA	155
3107	TSST1_NC002 758.2_334_3 57_F	TGCCAACATACTA GCGAAGGAACT	331	TSST1_NC00275 8.2_415_445_R	TCCCATGAACCTT AACTTTTAAAGGT AGTTC	332

[00107] As noted above, primer pair name codes for primer pairs listed in Table 1, crossreferenced to corresponding reference sequence, bioagent, and gene information are shown in Table
2. The primer name code typically represents the gene to which the given primer pair is targeted.
The primer names also include specific coordinates with respect to a reference sequence to which
the primer hybridizes. As exemplified above, this reference sequence is often defined by an
extraction of a section of sequence or defined by a GenBank gi number (indicated by extraction
coordinates in the primer pair name), or the corresponding complementary sequence of the
extraction, or, in cases when no extraction coordinates are listed, to the entire sequence of the
GenBank gi number. Gene abbreviations are shown in bold type in the "Gene Name" column of
Table 2.

[00108] Methods for PCR primer design are well known. One of skill in the art will understand that primer pairs configured to prime amplification of a double stranded sequence are configured and named using one strand of the double stranded sequence as a reference. The forward primer is the primer of the pair that comprises full or partial sequence identity to the one strand of the sequence being used as a reference. The reverse primer is the primer of the pair that comprises reverse complementarity to the one strand being of the sequence being used as a reference.

[00109] In one embodiment, the "plus" or "top" strand (the primary sequence as submitted to GenBank) of the nucleic acid to which the primers hybridize is used as a reference when designing primer pairs. In this case, the forward primer will comprise identity and the reverse primer will comprise reverse complementarity, to the sequence listed in GenBank for the reference sequence. In some embodiments, the primer pair is configured using the "minus" or "bottom" strand (reverse complement of the primary sequence as submitted to and listed in GenBank). In this case, the forward primer comprises sequence identity to the minus strand, and thus comprises reverse complementarity to the top strand, the sequence listed in GenBank. Similarly, in this case, the reverse primer comprises reverse complementarity to the minus strang, and thus comprises identity to the top strand.

[00110] Herein, when the primer is configured using the minus strand as a reference, the extraction sequence is preferably listed in a descending fashion in the primer name (as in the case of the coordinates 1674726-1674277 of the forward primer pair name AROE_NC003923-1674726-1674277_30_62_F). In this case, the forward primer comprises reverse complementarity to the sequence listed in GenBank for the reference gi number. Thus, in the case of this exemplary primer, the forward primer is configured to hybridize within nucleotides 1674697 and 1674665 of gi number 21281729, which is 30 (the first number in the hybridization coordinates 30-62) nucleotides in the reverse direction from the first coordinate (1674697) listed in the extraction sequence. The hybridization site and region of the reference sequence to which a primer in Table 1 hybridizes can be determined and verified with bioinformatics alignment tools as described below using the primer sequence and the reference gi number provided in Table 2.

[00111] To determine the exact primer hybridization coordinates of a given pair of primers on a given bioagent nucleic acid sequence and to determine the sequences, molecular masses and base compositions of an amplification product to be obtained upon amplification of nucleic acid of a known bioagent with known sequence information in the region of interest with a given pair of primers, one with ordinary skill in bioinformatics is capable of obtaining alignments of the primers of the present invention with the GenBank gi number of the relevant nucleic acid sequence of the known bioagent. For example, the reference sequence GenBank gi numbers (Table 2) provide the identities of the sequences which can be obtained from GenBank. Alignments can be done using a bioinformatics tool such as BLASTn provided to the public by NCBI (Bethesda, MD). Alternatively, a relevant GenBank sequence may be downloaded and imported into custom programmed or commercially available bioinformatics programs wherein the alignment can be carried out to determine the primer hybridization coordinates and the sequences, molecular masses and base compositions of the amplification product. For example, to obtain the hybridization coordinates of primer pair number 2095 (SEO ID NO.: 39; SEO ID NO.:125), First the forward primer (SEO ID NO: 39) is subjected to a BLASTn search on the publicly available NCBI BLAST website, "RefSeq Genomic" is chosen as the BLAST database since the gi numbers refer to genomic sequences. The BLAST query is then performed. Among the top results returned is a match to GenBank gi number 21281729 (Accession Number NC 003923). The result shown below. indicates that the forward primer hybridizes to positions 1530282...1530307 of the genomic sequence of Staphylococcus aureus subsp. aureus MW2 (represented by gi number 21281729).

```
Staphylococcus aureus subsp. aureus MW2, complete genome Length=2820462
```

Features in this part of subject sequence:
Panton-Valentine leukocidin chain F precursor

```
Score = 52.0 bits (26), Expect = 2e-05
Identities = 26/26 (100%), Gaps = 0/26 (0%)
Strand=Plus/Plus
```

[00112] The hybridization coordinates of the reverse primer (SEQ ID NO: 125) can be determined in a similar manner and thus, the bioagent identifying amplicon can be defined in terms of genomic coordinates. The query/subject arrangement of the result would be presented in Strand = Plus/Minus format because the reverse strand hybridizes to the reverse complement of the genomic sequence. The preceding sequence analyses are well known to one with ordinary skill in bioinformatics and thus, Table 2 contains sufficient information to determine the primer hybridization coordinates of any of the primers of Table 1 to the applicable reference sequences described therein.

Table 2: Primer Name Codes and Reference Sequences

Gene Name		Reference GenBank gi
1		33591275
		33591275
RNase P (ribonuclease 2)		
		53723370
RNase P (ribonuclease P)		16077068
		18308982
RNase P (ribonuclease ?)	Escherichia coli	16127994
	Rickettsia	15603881
RNase P (ribonuclease ?)	prowazekii	
	Staphylococcus	15922990
RNase P (ribonuclease P)	aureus	
RNase P (ribonuclease P)	Vibrio cholerae	15640032
icd (isocitrate dehydrogenase)	Ccxiella burnetii	29732244
	Acinetobacter	29732244
multi-locus IS1111A insertion element	baumannii	
	Rickettsia	40287451
ompA (outer membrane protein A)	prowazekii	
	Rickettsia	15603881
ompB (outer membrane protein E)	prowazekii	
gltA (citrate synthase)	Vibrio cholerae	15603881
	Francisella	15640032
toxR (transcription regulator toxR)	tularensis	
	Francise!la	56707187
dehydrogenase)	tularensis	
		56707187
	RNace P (ribonuclease 2) RNace P (ribonuclease 3) RNace P (ribonuclease 3)	Organism RNase P (ribonuclease ?) RNase P

Primer name code	Gene Name	Organism	Reference GenBank gi number
IPAH SCF	ipaH (invasion plasmid antigen)	Campylcbacter jejuni	30061571
HUPB_CJ	hupB (DNA-binding protein Hu-beta)	Coxiella burnetii	15791399
MUPR_X75439	mupR (mupriocin resistance gene;	Staphylococcus auseus	438226
PARC X95819	parC (topcisomerase IV)	Acinetobacter baumannii	1212748
SED_M2852_	sed (en-erotoxin D)	Staphylococcus aureus	1492109
SEJ_AF053140	sej (enterotoxin J)	Staphylococcus auzeus	3372540
AGR-III NCC03923	agr-III (accessory gene regulator-III)	Staphylococcus aureus	21281729
ARCC_NC003923	arcC (carbamate kinase)	Staphylococcus aureus	21281729
AROE_NC003923	aroE (shikimate 5-dehvdrcgenase	Staphylococcus aureus	21281729
BSA-A_NC003923	bsa-a (glutathione peroxidase)	Staphylococcus aureus	21281729
BSA-B_NCCC3923	bsa-b (epidermin bicsynthesis protein RpiB)	Staphylococcus aureus	21281729
GLPF_NC003923	glpF (qlycerol transporter)	Staphylococcus aureus	21281729
GMK_NC003923	gmk (guanylate kinase)	Staphylococcus aureus	21281729
MECT-R_NG003923	mecR1 (truncated methicillin resistance protein)	Staphylococcus aureus	21281729
PTA_NC003923	pta (phosphate acetyltransferase)	Staphylococcus aureus	21201729
PVLUK_NC003923	pvluk (Panton-Valentine leukocidin chain F precursor)	Staphylococcus aureus	21281729
SA442_NC003923	sa442 gene	Staphylococcus aureus	21281729
SEA_NC003923	sea (staphylococcal enterotoxin A precursor)	Staphylococcus aureus	21201729
SEC_NC003923	sec4 (enterotoxin type C precursor)	Staphylococcus aureus	21281729
TPI_NC003923	tpi (triosephosphate isomerase)	Staphylococcus aureus	21281729
YQI_NC003923	yqi (acetyl-CoA C-acetyltransferase homologue)	Staphylococcus aureus	21281729
AGR-II_NC002745	agr-II (accessory gene regulator-II)	Staphylococcus aureus	29165615
AGR-I_AJ617706	agr-I (accessory gene regulator-I)	Staphylococcus aureus	46019543
AGR-IV_AJ617711	agr-IV (accessory gene regulator-III)	Staphylococcus aureus	46019563
BLAZ_NC002952	blaZ (beta lactamase III)	Staphylococcus aureus	49482253
ERMA_NC002952	crmA (iRNA methylitansferase A)	Staphylococcus aureus	49482253
ERMB Y13600	ermB (rRNA methyltransferase B)	Staphylococcus aureus	49482253
SEA-SEE_NCC02952	sea (staphylococcal enterotoxin A precursor)	Staphylococcus sureus	49482253
SEA-SEE NCC02952	sea (staphylococcal enterotoxin A precursor)	Staphylococcus aureus	49482253
SEE_NC002952	sea (staphylococcal enterotoxin A procursor)	Staphylococcus aurous	49482253
SEH_NC002953	sch (staphylococcal enterotoxin H)	Staphylococcus aureus	49484912

Primer name code	Gene Name	Organism	Reference GenBank gi number
ERMC_NC005908	ermC (rRNA methyltransforase C)	Staphylococcus cureus	49489772
NUC NC002758	nuc (staphylococcal nuclease)	Staphylococcus cureus	15922990
SEB NC002758	seb (enterotoxin type B precursor)	Staphylococcus auseus	57634611
SEG NC002758	seg (staphylococcal enterotoxin G)	Staphylococcus aurous	57634611
SLI NC002758	sei (staphy occodal enterotoxin T)	Staphylococcus aureus	57634611
TSST_NC002758	tsst (toxic shock syndrome toxin-1)	Staphylococcus auseus	15922990
TUFB NC002758	tufB (Elongation factor Tu)	Staphylococcus aureus	15922990
188T1 NC002758.2	tsst (foxic shock syndrome Toxin=1)	Staphylococcus aureus	57634611

Example 2: Sample Preparation and PCR

[00113] Samples were processed to obtain bacterial genomic material using a Qiagen QIAamp Virus BioRobot MDx Kit (Valencia, CA 91355). Resulting genomic material was amplified using an MJ Thermocycler Dyad unit (BioRad laboratories, Inc., Hercules, CA 94547) and the amplicons were characterized on a Bruker Daltonics MicroTOF instrument (Billerica, MA 01821). The resulting molecular mass measurements were converted to base compositions and were queried into a database having base compositions indexed with primer pairs and bioagents.

[00114] All PCR reactions were assembled in 50 .micro.L reaction volumes in a 96-well microtiter plate format using a Packard MPII liquid handling robotic platform (Perkin Elmer, Bostan, MA 02118) and M.J. Dyad thermocyclers (BioRad, Inc., Hercules, CA 94547). The PCR reaction mixture consisted of 4 units of Amplitaq Gold, Ix buffer II (Applied Biosystems, Foster City, CA), 1.5 mM MgCl.sub.2, 0.4 M betaine, 800 .micro. M dNTP mixture and 250 nM of each primer. The following typical PCR conditions were used: 95.deg.C for 10 min followed by 8 cycles of 95.deg.C for 30 seconds, 48.deg.C for 30 seconds, and 72.deg.C 30 seconds with the 48.deg.C annealing temperature increasing 0.9.deg.C with each of the eight cycles. The PCR was then continued for 37 additional cycles of 95.deg.C for 15 seconds, 56.deg.C for 20 seconds, and 72.deg.C 20 seconds. Those ordinarily skilled in the art will understand PCR reactions.

Example 3: Solution Capture Purification of PCR Products for Mass Spectrometry with Ion Exchange Resin-Magnetic Beads

[00115] For solution capture of nucleic acids with ion exchange resin linked to magnetic beads, 25 micro.1 of a 2.5 mg/mL suspension of BioClone amine terminated supraparamagnetic beads (San Diego, CA 92126) were added to 25 to 50 .micro.1 of a PCR (or RT-PCR) reaction containing approximately 10 pM of an amplicon. The above suspension was mixed for approximately 5 minutes by vortexing or pipetting, after which the liquid was removed after using a magnetic separator. The beads containing bound PCR amplicon were then washed three times with 50mM ammonium bicarbonate/50% MeOH or 100mM ammonium bicarbonate/50% MeOH, followed by three more washes with 50% MeOH. The bound PCR amplicon was eluted with a solution of 25mM piperidine, 25mM midazole, 35% MeOH which included peptide calibration standards.

Example 4: Mass Spectrometry and Base Composition Analysis

100116] The ESI-FTICR mass spectrometer is based on a Bruker Daltonics (Billerica, MA) Apex II 70e electrospray ionization Fourier transform ion cyclotron resonance mass spectrometer that employs an actively shielded 7 Tesla superconducting magnet. The active shielding constrains the majority of the fringing magnetic field from the superconducting magnet to a relatively small volume. Thus, components that might be adversely affected by stray magnetic fields, such as CRT monitors, robotic components, and other electronics, can operate in close proximity to the FTICR spectrometer. All aspects of pulse sequence control and data acquisition were performed on a 600 MHz Pentium II data station running Bruker's Xmass software under Windows NT 4.0 operating system. Sample aliquots, typically 15 micro.l, were extracted directly from 96-well microtiter plates using a CTC HTS PAL autosampler (LEAP Technologies, Carrboro, NC) triggered by the FTICR data station. Samples were injected directly into a 10 micro.l sample loop integrated with a fluidics handling system that supplies the 100 micro.l /hr flow rate to the ESI source. Ions were formed via electrosoray ionization in a modified Analytica (Branford, CT) source employing an off

axis, grounded electrospray probe positioned approximately 1.5 cm from the metalized terminus of a glass desolvation capillary. The atmospheric pressure end of the glass capillary was biased at 6000 V relative to the ESI needle during data acquisition. A counter-current flow of dry N sub.2 was employed to assist in the desolvation process. Ions were accumulated in an external ion reservoir comprised of an rf-only hexapole, a skimmer cone, and an auxiliary gate electrode, prior to injection into the trapped ion cell where they were mass analyzed. Ionization duty cycles > 99% were achieved by simultaneously accumulating ions in the external ion reservoir during ion detection. Each detection event consisted of 1M data points digitized over 2.3 s. To improve the signal-to-noise ratio (S/N), 32 scans were co-added for a total data acquisition time of 74 s.

[00117] The ESI-TOF mass spectrometer is based on a Bruker Daltonics MicroTOF.sup.TM. Ions from the ESI source undergo orthogonal ion extraction and are focused in a reflectron prior to detection. The TOF and FTICR are equipped with the same automated sample handling and fluidics described above. Ions are formed in the standard MicroTOF.sup.TM ESI source that is equipped with the same off-axis sprayer and glass capillary as the FTICR ESI source. Consequently, source conditions were the same as those described above. External ion accumulation was also employed to improve ionization duty cycle during data acquisition. Each detection event on the TOF was comprised of 75,000 data points digitized over 75 .micro.s.

[00118] The sample delivery scheme allows sample aliquots to be rapidly injected into the electrospray source at high flow rate and subsequently be electrosprayed at a much lower flow rate for improved ESI sensitivity. Prior to injecting a sample, a bolus of buffer was injected at a high flow rate to rinse the transfer line and spray needle to avoid sample contamination/carryover. Following the rinse step, the autosampler injected the next sample and the flow rate was switched to low flow. Following a brief equilibration delay, data acquisition commenced. As spectra were coadded, the autosampler continued rinsing the syringe and picking up buffer to rinse the injector and sample transfer line. In general, two syringe rinses and one injector rinse were required to minimize sample carryover. During a routine screening protocol a new sample mixture was injected every 106 seconds. More recently a fast wash station for the syringe needle has been implemented which,

when combined with shorter acquisition times, facilitates the acquisition of mass spectra at a rate of just under one spectrum/minute.

[00119] Raw mass spectra were post-calibrated with an internal mass standard and deconvoluted to monoisotopic molecular masses. Unambiguous base compositions were derived from the exact mass measurements of the complementary single-stranded oligonucleotides. Quantitative results are obtained by comparing the peak heights with an internal PCR calibration standard present in every PCR well at 500 molecules per well. Calibration methods are commonly owned and disclosed in PCT pre-grant publication number WO 2005/094421, which is incorporated herein by reference in entirety.

Example 5: De Novo Determination of Base Composition of Amplicons using Molecular Mass Modified Deoxynucleotide Triphosphates.

[00120] Because the molecular masses of the four natural nucleobases have a relatively narrow molecular mass range (A = 313.058, G = 329.052, C = 289.046, T = 304.046, values in Daltons – See Table 3), a persistent source of ambiguity in assignment of base composition can occur as follows: two nucleic acid strands having different base composition may have a difference of about 1 Da when the base composition difference between the two strands is $G \leftrightarrow A$ (-15.994) combined with $C \leftrightarrow T$ (+15.000). For example, one 99-mer nucleic acid strand having a base composition of A.sub.27G.sub.30C.sub.21T.sub.21 has a theoretical molecular mass of 30779.058 while another 99-mer nucleic acid strand having a base composition of A.sub.26G.sub.31C.sub.22T.sub.20 has a theoretical molecular mass of 30780.052 is a molecular mass difference of only 0.994 Da. A 1 Da difference in molecular mass may be within the experimental error of a molecular nucleobases measurement and thus, the relatively narrow molecular mass range of the four natural nucleobases imposes an uncertainty factor in this type of situation. One method for removing this theoretical 1 Da uncertainty factor uses amplification of a nucleic acid with one mass-tagged nucleobase and three natural nucleobases.

[00121] Addition of significant mass to one of the 4 nucleobases (dNTPs) in an amplification reaction, or in the primers themselves, will result in a significant difference in mass of the resulting amplicon (greater than 1 Da) arising from ambiguities such as the G ↔ A combined with C ↔ T event (Table 3). Thus, the same the G ↔ A (-15.994) event combined with 5-lodo-C ↔ T (-110.900) event would result in a molecular mass difference of 126.894 Da. The molecular mass of the base composition $A_{27}G_{30}5$ -lodo- $C_{21}T_{21}$ (33422.958) compared with A.sub.26G.sub.315-lodo-C.sub.22T.sub.20, (33549.852) provides a theoretical molecular mass difference is +126.894. The experimental error of a molecular mass measurement is not significant with regard to this molecular mass difference. Furthermore, the only base composition consistent with a measured molecular mass of the 99-mer nucleic acid is A.sub.27G.sub.305-lodo-C.sub.21T.sub.21. In contrast, the analogous amplification without the mass tag has 18 possible base compositions.

Table 3: Molecular Masses of Natural Nucleobases and the Mass-Modified Nucleobase 5-Iodo-C and Molecular Mass Differences Resulting from Transitions

fucleobase	Molecular Mass	Transition	Δ Molecular Mass
A	313.058	A>T	-9.012
A	313.058	A>C	-24.C12
A	313.058	A>5-Ioqo-C	101.888
A	313.058	A>G	15.994
T	301.046	T>A	9.012
T	304.046	T>C	-15.000
T	304.046	T>5-Ioqo-C	110.900
T	304.046	T>G	25.006
С	289.046	C>A	24.012
С	289.046	C>T	15.000
С	289.046	C>G	40.006
5-Todo-C	414.946	5=Todo=C==>A	-101.888
5-Iodo-C	414.946	5-Iodo-C>T	-110.900
5-Todo-C	414.946	5-Todo-C>G	-85.894
G	329.052	G>A	-15.994
G	329.052	G==>T	-25.006
G	329.052	G>C	-40.006
G	329.052	G>5-1oco-C	85.894

[00122] Mass spectra of bioagent-identifying amplicons can be analyzed using a maximumlikelihood processor, such as is widely used in radar signal processing. This processor first makes maximum likelihood estimates of the input to the mass spectrometer for each primer by running matched filters for each base composition aggregate on the input data. This includes the response to a calibrant for each primer.

[00123] The algorithm emphasizes performance predictions culminating in probability-of-detection versus probability-of-false-alarm plots for conditions involving complex backgrounds of naturally occurring organisms and environmental contaminants. Matched filters consist of a priori expectations of signal values given the set of primers used for each of the bioagents. A genomic sequence database is used to define the mass base count matched filters. The database contains the sequences of known bacterial bioagents and includes threat organisms as well as benign background organisms. The latter is used to estimate and subtract the spectral signature produced by the background organisms. A maximum likelihood detection of known background organisms is implemented using matched filters and a running-sum estimate of the noise covariance. Background signal strengths are estimated and used along with the matched filters to form signatures which are then subtracted. The maximum likelihood process is applied to this "cleaned up" data in a similar manner employing matched filters for the organisms and a running-sum estimate of the noise-covariance for the cleaned up data.

[00124] The amplitudes of all base compositions of bioagent-identifying amplicons for each primer are calibrated and a final maximum likelihood amplitude estimate per organism is made based upon the multiple single primer estimates. Models of all system noise are factored into this two-stage maximum likelihood calculation. The processor reports the number of molecules of each base composition contained in the spectra. The quantity of amplicon corresponding to the appropriate primer set is reported as well as the quantities of primers remaining upon completion of the amplification reaction.

[00125] Base count blurring can be carried out as follows. Electronic PCR can be conducted on nucleotide sequences of the desired bioagents to obtain the different expected base counts that could

be obtained for each primer pair. See for example, Schuler, Genome Res. 7.541-50, 1997; or the e-PCR program available from National Center for Biotechnology Information (NCBI, NIH, Bethesda, MD 20894). One illustrative embodiment uses one or more spreadsheets from a workbook comprising a plurality of spreadsheets (e.g., Microsoft Excel). First in this example, there is a worksheet with a name similar to the workbook name; this worksheet contains the raw electronic PCR data. Second, there is a worksheet named "filtered bioagents base count" that contains bioagent name and base count; there is a separate record for each strain after removing sequences that are not identified with a genus and species and removing all sequences for bioagents with less than 10 strains. Third, there is a worksheet, "Sheet1" that contains the frequency of substitutions, insertions, or deletions for this primer pair. This data is generated by first creating a pivot table from the data in the "filtered bioagents base count" worksheet and then executing an Excel VBA macro. The macro creates a table of differences in base counts for bioagents of the same species, but different strains. One of ordinary skill in the art understands the additional pathways for obtaining similar table differences without undo experimentation.

[00126] Application of an exemplary script, involves the user defining a threshold that specifies the fraction of the strains that are represented by the reference set of base counts for each bioagent. The reference set of base counts for each bioagent may contain as many different base counts as are needed to meet or exceed the threshold. The set of reference base counts is defined by taking the most abundant strain's base type composition and adding it to the reference set and then the next most abundant strain's base type composition is added until the threshold is met or exceeded. The current set of data was obtained using a threshold of 55%, which was obtained empirically.

[00127] For each base count not included in the reference base count set for that bioagent, the script then proceeds to determine the manner in which the current base count differs from each of the base counts in the reference set. This difference may be represented as a combination of substitutions, Si=Xi, and insertions, Ii=Yi, or deletions, Di=Zi. If there is more than one reference base count, then the reported difference is chosen using rules that aim to minimize the number of changes and, in instances with the same number of changes, minimize the number of insertions or deletions. Therefore, the primary rule is to identify the difference with the minimum sum (Xi+Yi)

or (Xi+Zi), e.g., one insertion rather than two substitutions. If there are two or more differences with the minimum sum, then the one that will be reported is the one that contains the most substitutions.

[00128] Differences between a base count and a reference composition are categorized as one, two, or more substitutions, one, two, or more insertions, one, two, or more deletions, and combinations of substitutions and insertions or deletions. The different classes of nucleobase changes and their probabilities of occurrence have been delineated in U.S. Patent Application Publication No. 2004209260, which is incorporated herein by reference in entirety.

Example 6: Staphylococcus Bacterial Surveillance Panel.

[00129] The compositions and methods described herein are useful for screening a sample suspected of comprising one or more unknown bioagents to determine the identity of at least one of the bioagents. The compositions and methods provided are also useful for determining population genotype for a sample suspected of comprising a population of bioagents. In one embodiment, the population is a mixed population. The identification of the at least one bioagent or one or more genotypes is accomplished by generating base composition signatures using the methods provided herein for portions of genes shared by two or more members of the Staphylococcus genus. The base composition signatures generated using the methods provided are then compared to a database composition a plurality of base composition signatures that are indexed to primer pairs used in generating the base composition signatures and bioagents. The plurality of base composition signatures in the database is at least two, is more preferably at least 5, is more preferably still at least 14, is more preferably still at least 19, is more preferably still at least 25 and is more preferably still at least 19. The base composition signatures comprising this plurality identify at least one bioagent when that bioagent's measured and calculated base composition signature is queried against the plurality of base composition signatures comprised in the database.

Example 7 Identification of Drug Resistance Genes and Virulence Factors in Staphylococcus aureus

[00130] Three primer pair panels, each comprising eight primer pairs, were configured for identification of the *Staphylococcus aureus* species and for identification of drug resistance genes and virulence factors of *Staphylococcus aureus* bioagents. These panels are shown in Tables 4-6. The primer sequences in these panels can also be found in Table 1, and are cross-referenced in Tables 4-6 by primer pair numbers, primer pair names, and SEQ ID NOs.

Table 4: Panel of Primer Pairs for Identification of Drug Resistance Genes and Virulence
Factors in Staphylococcus aureus

Primer Pair No.	Forward Primer Name	Forward Primer (SEQ ID NO:)	Reverse Primer Name	Reverse Primer (SEQ ID NO:)	Target Gene
879	MECA Y14051 4507 4530 F	58	MECA Y14051 4555 4581 R	142	mecA
2056	MECI-R_NC003923-41798- 41609 33 60 F	62	MECI-R NC003923-41798- 41609 86 113 R	147	MecI-R
2081	ERMA_NC002952=55890= 56621 366 395 F	294	ERMA_NC002952=55890= 56621 438 465 R	295	ermA
2086	ERMC_NC005908-2004- 2738 85 116 F	35	ERMC_NC005908-2004- 2738 173 206 R	121	ermC
2095	PVLUK NC003923-1529595- 1531285 688 713 F	39	PVLUK NC003923-1529595- 1531285 775 804 R	125	Pv-luc
2249	TUFB_NC002758-615039- 616222 696 /25 F	47	TUFB_NC002758-615038- 616222 /93 820 R	132	tufB
2256	NUC NC002758-894288- 894974_316_345_F	55	NUC NC002758-894288- 894974_396_421_R	139	Nuc
2313	MUPR X75439 2486 2516 F	21	MUFR X/5439 2548 25/4 R	104	mupR

Table 5: Panel of Primer Pairs for Identification of Drug Resistance Genes and Virulence
Factors in Staphylococcus aureus

Primer Pair No.	Forward Primer Name	Forward Primer (SEQ ID NO:)	Reverse Primer Name	Reverse Primer (SEQ ID NO:)	Target Gene
879	MECA Y14051 4507 4530 F	58	MECA Y14051 4555 4581 R	142	mecA
2056	MECI-R_NC003923-41798- 41609 33 60 F	62	MECI-R_NC003923-41798- 41609 36 113 R	147	MecI-R
2081	ERMA_NC002952-55890- 56621 366 395 F	294	ERMA_NC002952-55890- 56621 438 465 R	295	crmA
2086	ERMC_NC005908=2004= 2738 85 116 F	35	ERMC NC005908-2004- 2738 173 206 R	121	ermc.
2095	PVLUK_NC003923=1529595= 1531285 688 713 F	39	PVLUK_NC003923=1529595= 1531285 775 804 R	125	Pv=luc
2249	TUFB_NC002758-615039- 616222 696 725 F	47	TUFB_NC002758-615038- 616222 793 320 R	132	tufB
2256	NUC NC002758-894288- 894974 316 345 F	55	NUC_NCO02758-894288- 894974 396 421 R	139	Nuc
3016	MUPR X75439 2432 2510 F	22	MILES X75439 2551 2573 R	106	munR

Table 6: Panel of Primer Pairs for Identification of Drug Resistance Genes and Virulence
Factors in Staphylococcus aureus

Primer Pair No.	Forward Primer Name	Forward Primer (SEQ ID NO:)	Reverse Primer Name	Reverse Primer (SEQ ID NO:)	Target Gene
879	MECA Y14051 4507 4530 F	58	MECA Y14051 4555 4581 R	142	mecA
2056	MECI-R_NC303923-41798- 41609 33 60 F	62	MECI-R NC003923-41798- 41609 36 113 R	147	MecI=R
2081	ERMA_NC002952-55890- 56621 366 395 E	294	ERMA_NC002952-55990- 56621 438 465 R	295	ermA
2086	ERMC_NC005908-2004- 2738 85 116 F	35	ERMC_NC005908-2004- 2738 173 206 R	121	esrmC
2095	PVLUK NC003923-1529595- 1531285 638 713 F	39	PVLUK NC003923-1529595- 1531285 775 804 R	125	Pv=1u<
2249	TUFB_NC002758-615039- 616222 696 725 E	47	TUFB_NC002758-615038- 616222 793 320 R	132	LufB
2256	NUC NC002758-894288- 894974 316 345 F	55	NUC NC302753-894288- 894974 396 421 R	139	Nuc
3106	TSST1 NC002/58.2- 2137509- 2138213 519 546 F	70	TSST1_NC002/58.2- 2137509-2133213_593- 620 R	155	tsst1

[00131] Primer pair numbers 2256 and 2249 are confirmation primers configured with the aim of high-level identification of Staphylococcus aureus. The nuc gene is a Staphylococcus aureus-specific marker gene. The tufB gene is a universal housekeeping gene but the bioagent identifying amplicon defined by primer pair number 2249 provides a unique base composition (A43 G28 C19 T35) which distinguishes Staphylococcus aureus from other members of the genus Staphylococcus.

[00132] High level methicillin resistance in a given strain of Staphylococcus aureus is indicated by bioagent identifying amplicons defined by primer pair numbers 879 and 2056. Analyses have indicated that primer pair number 879 is not expected to prime S. sciuri homolog or Enterococcus facculis faciem ampicillin-resistant PBP5 homologs.

[00133] Macrolide and erythromycin resistance in a given strain of *Staphylococcus aureus* is indicated by bioagent identifying amplicons defined by primer pair numbers 2081 and 2086.

[00134] Resistance to mupriocin in a given strain of *Staphylococcus aureus* is indicated by bioagent identifying amplicons defined by primer pair numbers 2313 and 3016.

[00135] In the above panels, virulence in a given strain of Staphylococcus aureus can be indicated by bioagent identifying amplicons defined by primer pair numbers 2095 and 3106. Primer pair number 2095 can identify both the pvl (lukS-PV) gene and the lukD gene which encodes a homologous enterotoxin. A bioagent identifying amplicon of the lukD gene defined by primer pair number 2095 has a six nucleobase length difference relative to the lukS-PV gene. Further, primer pair number 3106 is configured to generate amplicons within the tsst-1 gene, which encodes for shock syndrome toxin, which causes toxic shock syndrome (TSS).

[00136] A total of 32 blinded samples of different strains of Staphylococcus aureus were provided by the Center for Disease Control (CDC). Each sample was analyzed by PCR amplification with the first of these eight primer pair panels (shown in Table 4), followed by purification and measurement of molecular masses of the amplification products by mass spectrometry. Base compositions for the amplification products were calculated. The base compositions provide the information summarized above for each primer pair. The results are shown in Tables 7A and 7B.

Table 7A: Drug Resistance and Virulence Identified in Blinded Samples of Various Strains of Staphylococcus aureus with Primer Pair Nos. 2081, 2086, 2095 and 2256

Sample Index No.	Primer Pair No.	Primer Pair No.	Primer Pair No.	Primer Pair No.
	2081 (ermA)	2086 (ermC)	2095 (pv-luk)	2256 (nuc)
CDC0010	-	-	PVL=/lukDI	+
CDC0015	-	-	PVL+/lukD+	+
CDC0019	-	+	PVL-/lukD+	+
CDC0026	+	-	PVL-/lukD+	+
CDC0030	+	-	PVL-/lukD+	+
CDC004	-	-	PVL+/lukD+	+
CDC0014	-	+	PVL+/lukD+	+
CDC008	-	-	PVL-/lukD+	+
CDC001	+	-	PVL-/lukD+	+
CDC0022	+	-	PVL=/lukD+	+
CDC006	+	-	PVL-/lukD+	+
CDC007	-	-	PVL-/lukD+	+
CDCVRSA1	+	-	PVL=/lukD+	+
CDCVRSA2	+	+	PVL=/lukD+	+

CDC0011	+	-	PVL-/lukD+	+
CDC0012	_	-	PVL+/lukD-	+
CDC0021	+	-	PVL-/lukD+	+
CDC0023	÷ ·	-	PVL=/lukD+	+
CDC0025	+	-	PVL=/lukD+	+
CDC005	-	-	PVL-/lukD+	+
CDC0018	+	-	PVL+/lukD-	+
CDC002	-	-	PVL=/lukD+	+
CDC0028	+	-	PVL-/lukD+	+
CDC003	-	-	PVt=/lukb+	+
CDC0013	-	-	PVL+/lukD+	+
CDC.0016	-	-	PVT-/lukb+	+
CDC0027	+	-	PVL-/lukD:	+
CDC0029	-	-	PVL+/lukD+	+
CDC0020	-	+	PVL-/lukD+	+
CDC0024	-	-	PVL=/lukD+	+
CDC0031	-	-	PVL=/lukD+	+

Table 7B: Drug Resistance and Virulence Identified in Blinded Samples of Various Strains of Staphylococcus aureus with Primer Pair Nos. 2249, 879, 2056, and 2313

Sample	Primer Pair No. 2249	Primer Pair No.	Primer Pair No.	Primer Pair No.
Index No.	(tufB)	879 (mecA)	2056 (mecI-R)	2313 (mupR)
CDC0010	Staphylococcus aurous	+	+	-
CDC0C15	Staphylccoccus aureus	-	-	-
CDC0C19	Staphylococcus aureus	+	+	-
CDC0026	Staphylococcus aureus	+	+	-
CDC0C30	Staphylococcus aureus	+	+	-
CDC004	Staphylococcus aureus	+	+	-
CDC0C14	Staphylccoccus aureus	+	+	-
CDC008	Staphylococcus aureus	+	+	-
CDC 0 0 1.	Staphylococcus aureus	+	+	-
CDC0C22	Staphylococcus aureus	+	+	-
CDC006	Staphylococcus aureus	+	+	+
CDC007	Staphylcooccus aureus	+	+	-
CDCVRSA1	Staphylococcus aureus	+	+	-
CDCVRSA2	Staphylococcus aureus	+	+	-
CDC0C11	Staphylococcus aureus	-	-	-
CDC0012	Staphylococcus aureus	+	+	-
CDC0C21	Staphylococcus aureus	+	+	-
CDC0C23	Staphylococcus aureus	+	+	-
CDC0C25	Staphylcooccus aureus	+	+	-
CDC005	Staphylccoccus aureus	+	+	-

CDC0018	Staphylococcus aureus	+	+	-
CDC002	Staphylococcus aureus	+	+	-
GDG0C58	Staphylccoccus aureus	+	+	-
CDC003	Staphylococcus aureus	+	+	-
CDC0C13	Staphylccoccus aureus	+	+	-
CDC0C16	Staphylccoccus aureus	+	+	-
CDC0027	Staphylococcus aurous	+	+	-
CDC0029	Staphylccoccus aureus	+	+	-
CDC0020	Staphylcooccus aurous	-	-	-
CDC0024	Staphylogoggus aureus	+	+	-
CDC0031	Staphylococcus scleiferi	-	-	-

[00137] Upon un-blinding of the samples illustrated in Tables 7A and 7B is was noted that each of the PVL+ identifications agreed with PVL+ identified in the same samples by standard PCR assays. These results indicate that the panel of eight primer pairs is useful for identification of drug resistance and virulence sub-species characteristics for Staphylococcus aureus. Thus, it is expected that a kit comprising one or more of the members of the panels provided in Tables 4-6, and/or one or more other drug-resistance or virulence-identifying primer pairs provided here will be a useful embodiment.

Example 8: Selection and Use of Triangulation Genotyping Analysis Primer Pairs for Staphylococcus aureus

[00138] To combine the power of high-throughput mass spectrometric analysis of bioagent identifying amplicons with the sub-species characteristic resolving power provided by triangulation genotyping analysis, two panels of eight triangulation genotyping analysis primer pairs were selected. Each of the primer pairs in these panels is configured to produce bioagent identifying amplicons within one of six different housekeeping genes, which are listed in Tables 8 and 9. The primer sequences are found in Table 1 and are cross-referenced by the primer pair numbers, primer pair names and SEQ ID NOs listed in Tables 8 and 9.

Table 8: Primer Pairs for Triangulation Genotyping Analysis of Staphylococcus aureus

Primer Pair No.	Forward Primer Name	Forward Primer (SEQ ID NO:)	Reverse Primer Name	Reverse Primer (SEQ ID NO:)	Target Gene
2146	ARCC_NC003923-2725050- 2724595 131 161 F	72	ARCC_NC003923-2725050- 2724595 214 245 R	156	arcC
2149	AROE NC003923-1674726- 1674277 30 62 F	79	AROE NC003923=1674726= 1674277 155 181 R	166	aroE
2150	AROE NC003923-1674726- 1674277 204 232 F	76	ARGE NC003923-1674726- 1674277 308 335 R	162	aroE
2156	GMK_NC003923-1190906- 1191334 301 329 F	83	GMK NC003923-1190906- 1191334 403 432 R	170	gmk
2157	FTA NC003923-628885- 629355 237 263 F	87	PTA NC303923-628885- 629355 314 345 R	172	pta
2161	TPI_NC003923-830671- 831072 1 34 F	90	TPI_NC303923-830671- 831072 97 129 R	177	tpi
2163	YQI NC003923-378916- 379431 142 167 F	93	YQI NC303923-378916- 379431 259 284 R	180	yqi
2166	YQI_NC003923-378916- 3/9431_2/5_300_F	94	YQI_NC303923-378916- 3/9431_364_396_R	181	yqi

Table 9: Primer Pairs for Triangulation Genotyping Analysis of Staphylococcus aureus

Primer Pair No.	Forward Primer Name	Forward Primer (SEQ ID NO:)	Reverse Primer Name	Reverse Primer (SEQ ID NO:)	Target Gene
3025	ARCC_NC003923-2725050- 2724595 131 161 F	72	ARCC NC003923-2725050- 2724595 232 260 R	158	arcC
2149	AROE_NC003923-1674726- 16/42// 30 62 F	79	AROE_NC003923=1674726= 16/42// 155 181 R	166	aroE
2150	AROE_NC003923-1674726- 1674277 234 232 F	76	AROE_NC003923-1674726- 1674277 308 335 R	162	aroE
2156	GMK_NC003923-1190906- 1191334 301 329 F	83	GMK NC003923-1190906- 1191334 403 432 R	1/0	gmk
2157	FTA NC003923-628885- 629355 237 263 E	87	PTA NC303923-628885- 629355 314 345 K	172	pLa
2161	TPI NC003923-830671- 831072 1 34 F	90	TPI NC303923-830671- 831072 97 129 R	177	tpi
2163	YQ1_NC003923-378916- 379431 142 167 F	93	YQ1_NC303923-378916- 379431 259 284 R	180	yqi
2166	YOI_NC003923-378916- 379431_275_300_E	94	YQI_NC303923-378916- 379431_364_396_R	181	Adr

[00139] The samples that were analyzed for drug resistance and virulence in Example 7 were subjected to triangulation genotyping analysis with the first panel of primers listed above. The primer pairs of Table 8 were used to produce amplification products by PCR, which were subsequently purified and measured by mass spectrometry. Base compositions were calculated from the molecular masses and are shown in Tables 10A and 10B.

Table 10A: Triangulation Genotyping Analysis of Blinded Samples of Various Strains of Staphylococcus aureus with Primer Pair Nos. 2146, 2149, 2150 and 2156

Sample	i	Primer Pair No.	Primer Pair No.	Primer Pair No.	Primer Pair No.
Index No.	Strain	2146 (arcC)	2149 (aroE)	2150 (aroE)	2156 (gmk)
CDC0010	CO.7	A44 G24 C18 T29	A59 G21 C18 T51	M0 G36 C13 T43	A5C G30 C20 T32
CDC0315	607	A44 G24 C18 T29	A59 G24 C18 T51	A40 G36 C13 T43	A5C G30 C20 T32
CDC0319	COL	A44 G24 C18 T29	A59 G24 C10 T51	A40 G36 C13 T43	A5C G30 C20 T32
CDC0026	COT	A44 G24 C18 T29	A59 G24 C18 T51	A40 G36 C13 T43	A5C G30 C20 T32
CDC0030	CO.7	A44 G24 C18 T29	A59 G24 C18 T51	A40 G36 C13 T43	A5C G30 C20 T32
CDC804	602	A44 G24 C18 T29	A59 G24 C18 T51	A40 G36 C13 T43	A5C G30 C20 T32
CDC0014	CO.	A44 G24 C18 1/29	A59 G24 C18 T51	440 G36 C13 T43	ASC G30 C20 T32
CDC008	2277	A44 G24 C18 T29	A59 G24 C18 T51	A40 G36 C13 T43	A5C G30 C20 T32
CDC001	Mu50	A45 G23 C20 T27	A58 G24 C18 T52	A40 G36 C13 T43	A51 G29 C21 T31
CDC0022	Mu50	A45 G23 C20 T27	A58 G24 C18 T52	A40 G36 C13 T43	A51 G29 C21 T31
CDC006	MuS0	A45 G23 C20 T27	A58 G24 C18 T52	A40 G36 C13 T43	A51 G29 C21 T31
CDC0011	MRSA252	A45 G24 C18 T28	A58 G24 C19 T51	A41 G36 C12 T43	A51 G29 C21 T31
CDC0012	MRSA252	A45 G24 C18 T28	A58 G24 C19 T51	A41 G36 C12 T43	A51 G29 C21 T31
CDC0321	MRSA252	A45 G24 C18 T28	A58 G24 C19 T51	A41 G36 C12 T43	A51 G29 C21 T31
CDC0023	ST:110	A45 G24 C18 T28	A59 G24 C18 T51	A40 G36 C13 T43	A5C G30 C20 T32
CDC0325	ST:110	A45 G24 C18 T28	A59 G24 C18 T51	A40 G36 C13 T43	A5C G30 C20 T32
CECLUS	ST: 335	A44 G24 C18 T29	A59 G23 C19 T51	940 G36 C14 T42	A51 G29 C21 T31
CDC0018	ST:338	A44 G24 C18 T29	A59 G23 C19 T51	A40 G36 C14 T42	A51 G29 C21 T31
CDC002	ST:108	A46 G23 C20 T26	A58 G24 C19 T51	A42 G36 C12 T42	A51 G29 C20 T32
CDC0328	ST:108	A46 G23 C20 T26	A58 G21 C19 T51	A42 G36 C12 T42	A51 G29 C20 T32
CDC003	ST:107	A45 G23 C20 T27	A58 G24 C18 T52	A40 G36 C13 T43	A51 G29 C21 T31
CDC0013	87:12	ND	A59 G24 C18 T51	A40 G36 C13 T43	A51 G29 C21 T31
CDC0016	ST:120	A45 G23 C18 T29	A58 G24 C19 T51	A40 G37 C13 T42	A51 G29 C21 T31
CDC0327	ST:105	A45 G23 C20 T27	A58 G24 C18 T52	A40 G36 C13 T43	A51 G29 C21 T31
CDC0029	MSSA476	A45 G23 C20 T27	A58 G24 C19 T51	A40 G36 C13 T43	A5C G30 C20 T32
00000020	8"115	A44 G23 C2" 1'27	A59 G23 C1B 1152	A40 G36 C13 T43	A5C G30 C20 T32
CDC0324	ST:137	A45 G23 C20 T27	A57 G25 C19 T51	A40 G36 C13 T43	A51 G29 C22 T30
CDC0031	***	No product	No product	No product	No product

Table 10B: Triangulation Genotyping Analysis of Blinded Samples of Various Strains of Staphylococcus aureus with Primer Pair Nos. 2146, 2149, 2150 and 2156

Sample		Primer Pair No.	Primer Pair No.	Primer Pair No.	Primer Pair No.
Index No.	Strain	2157 (pta)	2161 (tpi)	2163 (yqi)	2166 (yqi)
CDC0010	607	A32 G25 C23 T29	A51 G28 C22 T28	A41 G37 C22 T43	A37 G30 C18 T37
CDC0015	007	A32 G25 C23 T29	A51 G28 C22 T28	A41 G37 C22 T43	A37 G30 C18 T37
CDC0019	CO2	A32 G25 C23 T29	A51 G28 C22 T28	A41 G37 C22 T43	A37 G30 C18 T37
CDC01926	COT-	A32 G25 G23 T29	A51 G28 C22 T28	841 G37 C22 T43	A37 G30 C'B T37
CDC0030	CO2	A32 G25 C23 T29	A51 G28 C22 T28	A41 G37 C22 T43	A37 G30 C18 T37
CDC004	002	A32 G25 C23 T29	A51 G28 C22 T28	A41 G37 C22 T43	A37 G30 C18 T37
CDC0314	COT	A32 G25 C23 T29	A51 G28 C22 T28	A41 G37 C22 T43	A37 G30 C18 T37
CDC008	unknown	A32 G25 C23 T29	A51 G28 C22 T28	A41 G37 C22 T43	A37 G30 C18 T37
CDC001	Mu50	A33 G25 C22 T29	A50 G28 C22 T29	A42 G36 C22 T43	A36 G31 C19 T35
CDC0022	Mu50	A33 G25 C22 T29	A50 G28 C22 T29	A42 G36 C22 T43	A36 G31 C19 T35
CDC006	Mu50	A33 G25 C22 T29	A50 G28 C22 T29	A42 G36 C22 T43	A36 G31 C19 T35
CDC0011	MRSA252	A32 G25 C23 T29	A50 G28 C22 T29	A42 G36 C22 T43	A37 G30 C18 T37
CDC0112	MRSA252	A32 G25 C23 T29	A50 (428 1722 1729	9.42 G36 C22 T43	A37 G30 C18 T37
CDC0321	MRSA252	A32 G25 C23 T29	A50 G28 C22 T29	A42 G36 C22 T43	A37 G30 C18 T37
CDC0023	ST:110	A32 G25 C23 T29	A51 G28 C22 T28	A41 G37 C22 T43	A37 G30 C18 T37
CDC0025	ST:110	A32 G25 C23 T29	A51 G28 C22 T28	A41 G37 C22 T43	A37 G30 C18 T37
CDC005	ST:338	A32 G25 C24 T28	A51 G27 C21 T30	A42 G36 C22 T43	A37 G30 C18 T37
CDC0010	ST:330	A32 G25 C24 T28	A51 G27 C21 T30	A42 G36 C22 T43	A37 G30 C18 T37
CDC002	ST:108	A33 G25 C23 T28	A50 G28 C22 T29	A42 G36 C22 T43	A37 G30 C18 T37
CDC0328	ST:108	A33 G25 C23 T28	A50 G28 C22 T29	A42 G36 C22 T43	A37 G30 C18 T37
CDC003	ST:107	A32 G25 C23 T29	A51 G28 C22 T28	A41 G37 C22 T43	A37 G50 C18 T37
(10010313	8":12	A32 G25 C23 129	A51 G28 (*22 1/28	442 G36 (*22 T43	A37 G30 C18 T37
CDC0016	ST:120	A32 G25 C24 T28	A50 G28 C21 T30	A42 G36 C22 T43	A37 G30 C18 T37

			A50 G28 C22 T29		
		A33 G25 C22 T29		A42 G36 C22 T43	
				A42 G36 C22 T43	
CDC0024	ST:137	A33 C25 C22 T29	A51 G28 C22 T28	A42 G36 C22 T43	A37 G30 C18 T37
CDC0031	***	A34 G25 C25 T25	A51 G27 C24 T27	No product	No product

[00140] Note: *** The sample CDC0031 was identified as Staphylococcus scleiferi as indicated in Example 7. Thus, the triangulation genotyping primers configured for Staphylococcus aureus would generally not be expected to prime and produce amplification products of this organism. Tables 10A and 10B indicate that amplification products are obtained for this organism only with primer pair numbers 2157 and 2161.

[00141] A total of thirteen different genotypes of Staphylococcus aureus were identified according to the unique combinations of base compositions across the eight different bioagent identifying amplicons obtained with the eight primer pairs. These results indicate that the eight primer pair panel is useful for analysis of unknown or newly emerging strains of Staphylococcus aureus, and thus it is expected that a kit comprising one or more of the members of the panels provided in Tables 8 and 9, and/or one or more other Staphylococcus aureus genotyping primer pairs provided herein, will be a useful embodiment.

Example 9: Survey of 326 Staphylococcus aureus Clinical Isolates Using Primers to Drug Resistance / Virulance and Triangulation Genotyping Analysis Primer Pairs

[00142] A total of 326 human clinical Staphylococcus aureus isolate samples were obtained from the Centers for Disease Control (CDC), Johns Hopkins University and University of Arizona. These samples were tested using a combination of 16 primer pairs comprising; the eight identification/resistance/virulence primer pairs listed in Table 4 and the eight genotyping primer pairs listed in Table 8. Virulence (PVL), antibiotic resistance (to Methicilin, Erythromycin and

Mupirocin), and strain type were determined for each of the 326 samples. Results are summarized in Table 11 and in Figure 2.

Table 11: Identification and Determination of Virulence and Drug Resistance of 326 Clinical Isolates using Staphylococcus aureus Primer Pair Panel

	Identificati	on	Virulence		Antibiotic Resistance					
				Met	hicillin	Erythromycin		Mupirocin		
# of Isolates	tufB	nuc	PVL	mecA	MccI-R	ermA	ermC	mupR		
81	S. aureus	+	-	+	+	+	-	-		
81	S. aureus	+	-	+	+	-	-	-		
34	S. aureus	+	-	+	+	+	-	-		
32	S. aureus	+	-	+	+	-	+	-		
30	S. aureus	+	+	+	+	-	-	-		
30	S. aureus	+	-	+	+	-	-	-		
10	S. aureus	+	-	+	+	-	+	-		
7	S. aureus	+	+	-		-	-	-		
3	S. aureus	+	-	+	+	+	-	+		

^{+:} presence of indicated gene/virulence/resistance; -: absence of indicated gene/virulence/resistance

[00143] As shown in Figure 2, Staphylococcus aureus strains USA 100, USA 300, USA 200/1100, and the extremely virulent USA 400 were identified among the 326 clinical isolate using the genotyping primer pairs used in this example. The genotyping data obtained using the methods provided here were consistent with data from by the agencies that provided the samples, obtained via pulse-field gel electrophoresis and sequencing. As illustrated in Table 11, tufB and nuc primer pairs confirmed that all 326 isolates belonged to the Staphylococcus aureus species. 37 samples exhibited virulence as identified by the presence of the PVL gene (as indicated by a "+"). Resistance to the indicated antibiotics ("+") was identified in a number of the samples. These drug resistance and virulence data were greater than 99% concordant with data from the agencies that provided the samples, obtained via standard phenotypic and PCR methods. Further, the data show that accurate and precise identification, genotype, virulence, and drug resistance information can be determined for a large group of clinical samples using a panel combining the identification,

characterization and genotyping primer pairs in Examples 7 and 8. This observation suggests that a kit comprising a combination of any of the primer pairs in the panel of primer pairs used in this example, or a combination of any of the other *Staphylococcus aureus* primer pairs provided herein configured to hybridize within the genes in this example will be a useful embodiment.

Example 10: Primer Pairs for Determining Resistance and Sensitivity to Quinolones

[00144] Table 12 illustrates four primer pairs that were configured to determine quinolone resistance or sensitivity of Staphylococcus aureus bioagents. The primers of these pairs were configured to hybridize within regions of the Staphylococcus aureus gyrA gene. Sequences for these primers can be found in Table 1, and the primers are cross-referenced by primer name and SEQ ID NO in Table 12.

Table 12: Primer Pairs for Identification of Ouinolone Resistance in Staphylococcus aureus

Primer Pair Number	Forward Primer Name	Forward Primer SEQ ID NO.	Reverse Primer Name	Reverse Primer SEQ ID NO.
2738	GYRA_NC002 953-7005- 9668_166_195_ F	2	GYRA_NC002 953-7005- 9668_265_287 _R	5
2739	GYRA_NC002 953-7005- 9668_221_249_ F	3	GYRA_NC002 953-7005- 9668_316_343 _R	6
2740	GYRA_NC002 953-7005- 9668_221_249_ F	3	GYRA_NC002 953-7005- 9668_253_283 _R	7
2741	GYRA_NC002 953-7005- 9668_234_261_ F	4	GYRA_NC002 953-7005- 9668_265_287 _R	5

[00145] Each of the primer pairs listed in Table 12 is configured to generate an amplicon within at least a portion of the QRDR region of the gyrA gene (SEQ ID NO.:10), which confers quinolone resistance or sensitivity. The QRDR comprises the position of a drug resistance-conferring SNP of

the gyrA gene sequence, comprising a change of a single "C" nucleobase to a "T" nucleobase that results in a leucine instead of a serine at amino acid of the gyrase A protein. In the case of the reference sequence used to configure the primer pairs of Table 12, the SNP is located at position 251 of the extraction sequence ((coordinates 7005-9668) SEQ ID NO.: 8), which is the gyrA gene, from GenBank gi number 49484912. Forward primers in Table 12 are configured to comprise sequence identity within SEQ ID NO.: 11, a region of GenBank gi number 49484912. The reverse primers in Table 12 are configured to comprise reverse complementarity within SEQ ID NO.: 12, another region of GenBank gi number 49484912. The gyrA primer pairs provided in Table 12, when used in the methods provided herein, can detect a single nucleotide change at this SNP position, and are thus able to determine the drug resistant/sensitive genotype for the gyrA gene for a given Staphylococcus aureus bioagent.

Example 11: Characterizing Staphylococcus aureus in a Patient Sample Using Quinolone Resistant Primer Pairs and Other Staphylococcus aureus Primer Pairs

[00146] Population genotypes for mixed populations of bioagents can be identified with high sensitivity by PCR-ESI/MS because amplified bioagent nucleic acids having different base compositions appear in different positions in the mass spectrum. The dynamic range for mixed PCR-ESI/MS detections has previously been determined to be approximately 100:1(Hofstadler, S. A. et al., Inter. J. Mass Spectrom. (2005) 242, 23), which allows for detection of genotype variants with as low as 1% abundance in a mixed population. This detection using PCR-ESI/MS surveillance does not require secondary testing.

[00147] A wound sample from a patient infected with Staphylococcus aureus was analyzed directly by the methods provided herein using a panel of 17 primer pairs comprising: the eight identification/resistance/virulence primer pairs listed in Table 4, the eight genotyping primer pairs listed in Table 8, and the quinolone resistance determining primer pair (number 2740, SEQ ID NO: 3:SEO ID NO:7) listed in Table 12.

[00148] The sample was analyzed directly as described above in the previous examples using the primer pairs of Table 4, 8, and 12 (listed along the top of Table 13) in the methods provided herein. Further, a portion of the sample was cultured on an agar plate over a period of 2 days for further testing. Following the two-day culture, 9 colonies were picked and nucleic acids there from analyzed by the 17 primer pairs described above using the methods provided herein. The results are summarized in Table 13 and Figure 3.

Table 13 Analysis of Patient Sample Comprising Mixed Population of Staphylococcus aureus Bioagents: Identification of Quinolone Resistant and Sensitive Genotypes

	I	D	Viru	lence		Antibiotic Resistance					
					Meth	Methicillin Erythromycin		Mupirocin	Quinolone		
	pp # 2249	pp #2256	pp #2095	pp #2095	6£8# dd	pp #2056	pp #2081	pp #2086	pp #2313	pp#2740	panel of Table 8
	tuf B	nuc	luk D	PV L	mecA	MecI- R	ermA	ermC	mupR	gyrA	
Wound	SA	+	+	+	+	+	-	-	-	75% - 25%+	USA300
Colony 1	SA	+	+	+	+	+	-	-	-	-	USA300
Colony 2	SA	+	+	+	+	+	-	-	-	-	USA300
Colony 3	SA	+	+	+	+	+	-	-	-	+	USA300
Colony 4	SA	+	+	+	+	+	-	-	-	-	USA300
Colony 5	SA	+	+	+	+	+	-	-	-	-	USA300
Colony 6	SA	+	+	+	+	+	-	-	-	-	USA300
Colony 7	SA	+	+	+	+	+	-	-	-	-	USA300
Colony 8	SA	+	+	+	+	+	-	-	-	+	USA300
Colony 9	SA	+	+	+	+	+	-	-	-	-	USA300

ID: Identification; pp#: primer pair number; SA: Staphylococcus aureus; +: presence of indicated gene/virulence/resistance; -: absence of indicated gene/virulence/resistance

[00149] As shown in Table 13, the wound sample, and all colonies grown from that sample were determined to comprise one or more bioagents, identified by the methods provided here as Strain USA300 of MRSA Staphylococcus aureus. These one or more bioagents comprised in all samples

were also determined to be viurulent (pvl, lukD), methicillin resistant (mecA, mecI-R), and sensitive to erythromycin and mupirocin (ermA, ermC, mupR).

[00150] However, use of primer pair # 2740, which is configured to generate amplicons within the gyrA gene, identified a mixed population of bioagents in the patient sample, with more than one distinguishable genotype for the gyrA gene. Figure 3 shows a mass spectrum for the sample generated using primer pair number 2740. The two peak groupings represent the forward and reverse strands of the amplicon. Two different base compositions for amplicons generated by the primer pair were identified in the sample, evidenced by the double peaks shown for each strand. These double peaks (and base compositions determined therefrom) indicate that two genotypes, differing only by a single nucleotide at a SNP position in gyrA, were present in the patient sample. One genotype, comprising a C at the SNP of the gyrA gene, confering quinolone sensitivity, resulted in an amplicon with the base composition A.sub.19 G.sub.13 C.sub.11 T.sub.20. The other, comprising a T at the SNP position, confering quinolone resistance, resulted in an amplicon with the basee composition: A.sub.19 G.sub.13 C.sub.10 T.sub.21. As shown in the spectrum, the lower abundance genotype was present at approximately 25% of the population. This result is also indicated in Table 13, which lists the population genotype for the gyrA gene (Ouinolone column). which comprises both quinolone resistant and quinolone sensitive genoytpes at 25 and 75% respectively.

[00151] Further, Table 13 shows that two of the nine colonies (colony 3 and 8) screened in this example were found to comprise quinolone resistance, while the other six colonies comprised quinolone sensitivity, supporting the finding that the double peaks in the spectrumfor the wound sample represent a mixed population with two distinguishable genotypes. A spectrum and a base composition for an example of each type of colony is also shown in Figure 3.

[00152] Thus, the primer pairs and methods used in this example identified a mixed population of Staphylococcus aureus bioagents in a patient sample, and identified the population genotype for this mixed population. The methods and primer pairs provided herein will likely be useful in identifying population genotypes, emerging genotypes, and emerging populations of bioagents. A kit

comprising a combination of any of the primer pairs used in this example or other gyrA primer pairs provided herein will likely be a useful embodiment.

Example 12: Periodic Analysis of Population Genotypes in a Sample over time

Io0153] A sample, obtained from a patient or other sample source will be monitored over time using the primer pairs provided herein configured to identify quinolone resistant or sensitive genotypes. In this example, nucleic acids from the sample, obtained from a patient or other source suspected of comprising one or more bioagents, will be amplified using one or more of the primer pairs from Table 12, from each of any Staphylococcus aureus bioagents comprised in the sample. A base composition and/or molecular mass obtained using the methods provided herein will be compared to a database comprising molecular masses and/or base compositions, each indexed to the primer pair used and a bioagent genotype. Thus, a population genotype will be identified for the gyrA gene that will indicate the presence or absence of quinolone resistant and/or sensitive Staphylococcus aureus bioagents in the sample source. Optionally, one or more additional primer pairs will be used, such as any of the primer pairs from Tables 4-6 and 8-9 will be used to determine other characteristics of the bioagents in the sample.

[00154] An antibiotic regimen tailored to the identified genotype or genotypes will then be administered to the sample source. If the population comprises only the quinolone sensitive genotype, the antibiotic regimen may comprise a quinolone. If at least a percentage of the bioagents in the population of bioagents in the sample source comprises the quinolone resistant genotype, the antibiotic regimen will comprise an antibiotic for treating quinolone resistant bacteria. Periodically, samples will be subsequently obtained from the source, and the method repeated to monitor for emerging genotypes. Following each periodic repeat of the method, it will be determined whether there is an emerging genotype in the population of bioagents in the sample. If, after the initial identification, quinolones are being used in the antibiotic regimen tailored to treat the sample source and an emerging quinolone resistant genotype is identified during the periodic testing, the regimen will be modified to treat quinolone resistant bacteria. This modification will comprise addition of an antibiotic for treating quinolone resistant bacteria, and may further comprise discontinuation of

treatment with quinolones. In one embodiment, a combination of quinolones and an antibiotic to treat quinolone resistant bacteria may be used.

[00155] Various modifications to the description herein will be apparent to those skilled in the art from the foregoing description. Such modifications fall within the spirit and scope of the current invention and appended claims. Each reference (including, but not limited to, journal articles, U.S. and non-U.S. patents, patent application publications, international patent application publications, gene bank accession numbers, internet web sites, and the like) cited in the present application is incorporated herein by reference in its entirety.

WHAT IS CLAIMED IS:

We claim:

- 1. A method for identifying a population genotype comprising the steps of:
 - (a) obtaining a sample suspected of comprising a population of bioagents;
- (b) amplifying a nucleic acid from each of two or more bioagents from said population of bioagents in said sample using a primer pair that is configured to generate an amplicon from within a region defined by SEQ ID NO: 10, thereby generating amplicons from said nucleic acids;
- (c) determining a molecular mass measurement for each of said amplicons using a mass spectrometer;
 - (d) calculating a base composition from each molecular mass measurement; and
- identifying a population genotype for said population of bioagents by comparing each
 of said base compositions calculated in step (d) to a database of base compositions indexed to the
 primer pair of step (b) and a known bioagent genotype.
- 2. The method of claim 1 wherein said primer pair further comprises a forward member that is 20 to 35 nucleobases in length and comprises at least 80% identity to a first portion of SEQ ID NO: 10 and a reverse member that is 20 to 35 nucleobases in length and comprises at least 80% reverse complementarity to a second portion of SEQ ID NO: 10.
- 3. The method of claim 2 wherein said forward member comprises at least 90% identity to said first portion of SEQ ID NO: 10.
- The method of claim 2 wherein said forward member comprises at least 95% identity to said first portion of SEQ ID NO: 10.
- 5. The method of claim 2 wherein said forward member comprises at least 97% identity to said first portion of SEQ ID NO: 10.

 The method of claim 2 wherein said forward primer pair member comprises SEQ ID NO: 2 with 0-8 nucleobase deletions, additions and/or substitutions.

- The method of claim 2 wherein said forward primer pair member comprises SEQ ID NO: 3
 with 0-8 nucleobase deletions, additions and/or substitutions.
- The method of claim 2 wherein said forward primer pair member comprises SEQ ID NO: 4 with 0-8 nucleobase deletions, additions and/or substitutions.
- The method of claim 2 wherein said reverse member comprises at least 90% reverse complementarity to said second portion of SEQ ID NO: 10.
- The method of claim 2 wherein said reverse member comprises at least 95% reverse complementarity to said second portion of SEQ ID NO: 10.
- 11. The method of claim 2 wherein said reverse member comprises at least 97% reverse complementarity to said second portion of SEO ID NO: 10.
- 12. The method of claim 2 wherein said reverse primer pair member comprises SEQ ID NO: 5 with 0-6 nucleobase deletions, additions and/or substitutions.
- 13. The method of claim 2 wherein said reverse primer pair member comprises SEQ ID NO: 6 with 0-8 nucleobase deletions, additions and/or substitutions.
- 14. The method of claim 2 wherein said reverse primer pair member comprises SEQ ID NO: 7 with 0-9 nucleobase deletions, additions and/or substitutions.
- 15. The method of claim 1 wherein either or both of said primer members comprises at least one modified nucleobase

The method of claim 15 wherein said modified nucleobase is a mass modified nucleobase.

- The method of claim 16 wherein said modified nucleobase is 5-Iodo-C.
- The method of claim 15 wherein said modified nucleobase is a universal nucleobase.
- The method of claim 18 wherein said modified nucleobase is inosine.
- The method of claim 1 wherein either or both of said primer members comprise a nontemplated 5' T-residue.
- 21. The method of claim 1 wherein said population of bioagents comprises at least two bacteria belonging to the *Staphylococcus* genus.
- 22. The method of claim 21 wherein at least one of said bacteria is resistant to quinolone antimicrobial therapy.
- 23. The method of claim 21 wherein at least one of said bacteria is resistant to quinolone antimicrobial therapy and at least one of said bacteria is sensitive to quinolone antimicrobial therapy.
- 24. The method of claim 1 wherein said population of bioagents comprises at least two bacteria belonging to the Staphylococcus aureus species.
- 25. The method of claim 24 wherein at least one of said bacteria is resistant to quinolone antimicrobial therapy.
- 26. The method of claim 24 wherein at least one of said bacteria is resistant to quinolone antimicrobial therapy and at least one of said bacteria is sensitive to quinolone antimicrobial therapy.

27. The method of claim 1 wherein an antibiotic regimen tailored to treat the identified genotypes for the population of bioagents is delivered to the sample source.

- 28. The method of claim 1 wherein steps (a) to (e) are periodically repeated.
- 29. A method of reducing a population of bacteria in a person needing such a treatment comprising the steps of:
- (a) obtaining from a person a sample suspected of comprising a population of bacterial bioagents;
- (b) amplifying a nucleic acid from each of two or more bacterial bioagents in said sample using a primer pair that is configured to generate an amplicon from within a region of defined by SEO ID NO: 10, thereby generating amplicons from said nucleic acids:
- (c) determining a molecular mass measurement for each of said amplicons using a mass spectrometer;
 - (d) calculating a base composition from each molecular mass measurement;
- (e) identifying a population genotype for said population of bioagents by comparing each of said base compositions calculated in step (d) to a database of base compositions indexed to the primer pair of step (b) and a known bioagent genotype; and
- (f) administering to a person in need thereof an antibiotic regimen tailored to treat the identified genotypes for the population of bacterial bioagents.
- 30. The method of claim 29 wherein said primer pair further comprises a forward member that is 20 to 35 nucleobases in length and comprises at least 80% identity to a first portion of SEQ ID NO: 10 and a reverse member that is 20 to 35 nucleobases in length and comprises at least 80% reverse complementarity to a second portion of SEQ ID NO: 10.
- 31. The method of claim 30 wherein said forward member comprises at least 90% identity to said first portion of SEQ ID NO: 10.

 The method of claim 30 wherein said forward member comprises at least 95% identity to said first portion of SEQ ID NO: 10.

- The method of claim 30 wherein said forward member comprises at least 97% identity to said first portion of SEO ID NO: 10.
- 34. The method of claim 30 wherein said forward primer pair member comprises SEQ ID NO: 2 with 0-8 nucleobase deletions, additions and/or substitutions.
- 35. The method of claim 30 wherein said forward primer pair member comprises SEQ ID NO: 3 with 0-8 nucleobase deletions, additions and/or substitutions.
- 36. The method of claim 30 wherein said forward primer pair member comprises SEQ ID NO: 4 with 0-8 nucleobase deletions, additions and/or substitutions.
- 37. The method of claim 30 wherein said reverse member comprises at least 90% reverse complementarity to said second portion of SEQ ID NO: 10.
- 38. The method of claim 30 wherein said reverse member comprises at least 95% reverse complementarity to said second portion of SEO ID NO: 10.
- The method of claim 30 wherein said reverse member comprises at least 97% reverse complementarity to said second portion of SEQ ID NO: 10.
- 40. The method of claim 30 wherein said reverse primer pair member comprises SEQ ID NO: 5 with 0-6 nucleobase deletions, additions and/or substitutions.
- 41. The method of claim 30 wherein said reverse primer pair member comprises SEQ ID NO: 6 with 0-8 nucleobase deletions, additions and/or substitutions.

The method of claim 30 wherein said reverse primer pair member comprises SEQ ID NO. 7
 with 0-9 nucleobase deletions, additions and/or substitutions.

- The method of claim 30 wherein either or both of said primer members comprises at least one modified nucleobase
- 44. The method of claim 43 wherein said modified nucleobase is a mass modified nucleobase.
- 45 The method of claim 44 wherein said modified nucleobase is 5-Iodo-C.
- 46 The method of claim 43 wherein said modified nucleobase is a universal nucleobase.
- The method of claim 46 wherein said modified nucleobase is inosine.
- 48. The method of claim 29 wherein either or both of said primer members comprise a non-templated 5' T-residue.
- 49. The method of claim 29 wherein said population of bacterial bioagents comprises at least two bacteria belonging to the *Staphylococcus* genus.
- 50. The method of claim 49 wherein at least one of said bacteria is resistant to quinolone antimicrobial therapy.
- 51. The method of claim 49 wherein at least one of said bacteria is resistant to quinolone antimicrobial therapy and at least one of said bacteria is sensitive to quinolone antimicrobial therapy.
- 52. The method of claim 29 wherein said population of bacterial bioagents comprises at least two bacteria belonging to the *Staphylococcus aureus* species.

53. The method of claim 52 wherein at least one of said bacteria is resistant to quinolone antimicrobial therapy.

- 54. The method of claim 52 wherein at least one of said bacteria is resistant to quinolone antimicrobial therapy and at least one of said bacteria is sensitive to quinolone antimicrobial therapy.
- 55. The method of claim 29 wherein steps (a) to (e) are periodically repeated.
- 56. The method of claim 55 wherein an emerging genotype is identified in step (e) of one or more of said periodic repeats, further comprising modifying said antibiotic regimen to treat said emerging genotype.
- 57. The method of claim 29 wherein said antibiotic regimen comprises an antibiotic for treating quinolone resistant bacteria and an antibiotic for treating quinolone sensitive bacteria.
- 58. A composition of matter comprising a purified oligonucleotide primer pair wherein each primer member of said primer pair is 20 to 35 nucleobases in length and wherein the forward primer comprises at least 80% identity with a first portion of SEQ ID NO: 10 and the reverse primer comprises at least 80% reverse complementarity with a second portion of SEQ ID NO: 10.
- 59. The composition of claim 58 wherein the forward member comprises at least 90% identity to said first portion of SEQ ID NO: 10.
- The composition of claim 58 wherein the forward member comprises at least 95% identity to said first portion of SEO ID NO: 10.
- The composition of claim 58 wherein the forward member comprises at least 97% identity to said first portion of SEQ ID NO: 10.

The composition of claim 58 wherein the forward primer pair member comprises SEQ ID
 vith 0-8 nucleobase deletions, additions and/or substitutions.

- 63. The composition of claim 58 wherein the forward primer pair member comprises SEQ ID NO: 3 with 0-8 nucleobase deletions, additions and/or substitutions.
- 64. The composition of claim 58 wherein the forward primer pair member comprises SEQ ID NO: 4 with 0-8 nucleobase deletions. additions and/or substitutions.
- 65. The composition of claim 58 wherein the forward primer pair member comprises at least 80% identity with a portion of SEQ ID NO: 11.
- 66. The composition of claim 58 wherein the reverse member comprises at least 90% reverse complementarity to said second portion of SEQ ID NO: 10.
- The composition of claim 58 wherein the reverse member comprises at least 95% reverse complementarity to said second portion of SEQ ID NO: 10.
- The composition of claim 58 wherein the reverse member comprises at least 97% reverse complementarity to said second portion of SEQ ID NO: 10.
- The composition of claim 58 wherein the reverse primer pair member comprises SEQ ID
 NO: 5 with 0-6 nucleobase deletions, additions and/or substitutions.
- The composition of claim 58 wherein the reverse primer pair member comprises SEQ ID
 NO: 6 with 0-8 nucleobase deletions, additions and/or substitutions.
- 71. The composition of claim 58 wherein the reverse primer pair member comprises SEQ ID NO: 7 with 0-9 nucleobase deletions, additions and/or substitutions.

72. The composition of claim 58 wherein the reverse primer pair member comprises at least 80% reverse complementarity with a portion of SEQ ID NO: 12.

- 73. The composition of claim 58 wherein either or both of the primer members comprises at least one modified nucleobase.
- 74. The composition of claim 73 wherein the modified nucleobase is a mass modified nucleobase.
- 75. The composition of claim 74 wherein the modified nucleobase is 5-Iodo-C.
- The composition of claim 73 wherein the modified nucleobase is a universal nucleobase.
- 77. The composition of claim 76 wherein the modified nucleobase is inosine.
- 78. The composition of claim 58 wherein either or both of the primer members comprise a nontemplated 5' T-residue.
- 79. The composition of claim 58 wherein said primer pair is configured to generate an amplicon of between about 45 and about 192 nucleobases in length comprising a region of SEQ ID NO: 10.

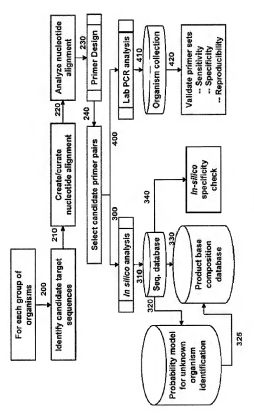


Figure 1

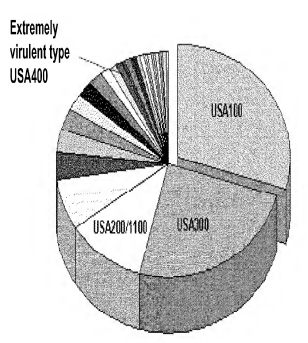


Figure 2

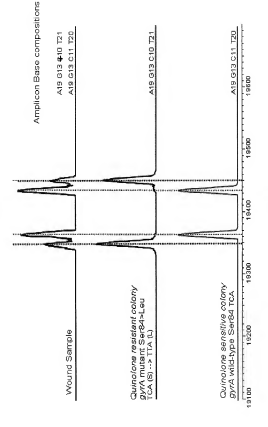


Figure 3

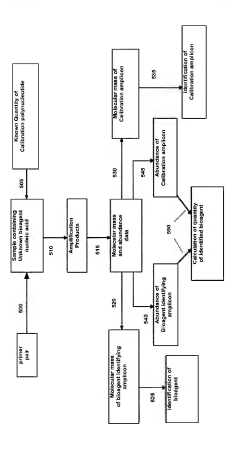


Figure 4